# OPTIMAL DESIGN OF A BIOGAS POWERED Libr-H<sub>2</sub>O VAPOUR ABSORPTION SYSTEM

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ME me/1985/m 1086-51130



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FEBRUARY, 1985

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A Thesis Submitted
in Partial Fulfilment of the Requirements
for the Degree of

MASTER OF TECHNOLOGY

By
MOHD. ALTAMUSH SIDDIQUI

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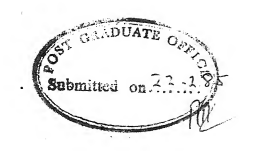
DEPARTMENT OF MECHANICAL ENGINEERING
INDIAN INSTITUTE OF TECHNOLOGY KANPUR
FEBRUARY, 1985

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#### CERTIFICATE

This is to certify that the thesis entitled "Optimal Design of a Biogas Powered LiBr-H, O Vapour Absorption System" by Mohd. Altamush Siddiqui is a record of work carried out under our supervision and has not been submitted elsewhere for a degree.

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This thesis has been approved for the award of the Degree of Master of Technology (M.Tech.) in accordance with the regulations of the Indian Institute of Technology Nangur Dated. 70

#### **ACKNOWL EDGEMENT**

I express my whole hearted gratitude to

Dr. Manohar Prasad and Dr. B. Sahay for their valuable

guidance and critical appraisal throughout the work.

Their constant inspiration and encouragement helped in

carrying out the experimental setup.

Thanks to Dr. D.P. Rao (Chemical Engg. Department) and Dr. K.K. Saxena (Mechanical Engg. Department) for their suggestions in the design of the components and the experimental setup of the system.

Thanks to Mr. P.N. Misra for his assistance in the assembly of the system. Thanks to the technical staffs of the Central Workshop, TA 203 Lab. and Glass Blowing, etc. Thanks to Mr. Sharma and his staff for their patience in fabricating the glass instruments for the measurement of pressure, temperature, etc.

Thanks to all my friends and well wishers for their kind help.

Thanks to Mr. D.P. Saini and Mr. Ayodhya Prasad for their patience in typing and Cyclostyling.

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# NOMENCLATURES

A	: cross-sectional area, m <sup>2</sup>
<sup>A</sup> f	: fin surface area, m <sup>2</sup>
Ao	: outer cross sectional area, m <sup>2</sup>
${}^{\rm A}{\tt i}$	: inner cross sectional area, m <sup>2</sup>
C <sub>1</sub>	: capital cost of biogas plant, Rupees
C <sub>2</sub>	: yearly cost of biogas plant, Rupees/year
$C_{\mathrm{T}}$	: total yearly running cost of biogas plant,
	Rupees/year
C <sub>p</sub>	: specific heat, kJ/kg-°C
C <sub>pw</sub>	: specific heat of water, kJ/kg-°C
C <sub>pL</sub> C <sub>sf</sub>	: specific heat of LiBr-H2O, solution, kJ/kg-°C
C <sub>s</sub>	: constant described by the liqd. surface
<b>L</b> ,	combination
COP	: coeff. of performance of refrigerating system
đ	: tube diameter, m
D	: Pipe, cylinder diameter, m
D	: bubble diameter, im (defined by Eq. 4.35)
FF	: finning factor, defined by Eq. (4.9)
F <sub>1-2</sub>	: shape factor from body 1 to body 2
G	: mass flux, kg/m <sup>2</sup> -h
h	: heat transfer coefficient, kJ/m <sup>2</sup> -h-°C
h.	: latent heat of vaporization/condensation

```
hiv
          : effective latent heat of vaporization/condensation
             allowing for the effect of superheat, kJ/m2-h-°C
h<sub>red</sub>
          : reduced heat transfer coefficient defined by
            Eq. (4.10)
          : inside heat transfer coeff. kJ/m2-h-°C
h;
h<sub>o</sub>
          : outside heat transfer coeff. kJ/m2-h-°C
          : fin side heat transfer coeff. kJ/m2-h-°C
h
^{\rm h}{\rm r_{\rm g}}
          : radiative heat transfer coeff. in the
            generator, kJ/m2-h-°C
hr
          : energy release due to respiration per tonne
            mass per unit time
hc
          : convective heat transfer coefficient kJ/m2-h-°C
          : thermal conductivity, kJ/m-h-°C
k
          : characteristic lengthfor fin, m
          : length of tubing, piping of the respective
L
suffix
            system components, m
L
          : life of the biogas plant, years
          : mass flow at any section per unit mass flow through evaporator (kg/kg)
          : mass flow rate, kg/h (suffixes indicate the state points)
          : number of fins per meter length
n
          : heat transfer rate per unit length of the
q
            tube, kJ/h
Ō
          : heat transfer rate per TR, kJ/h
          : total heat transfer rate, kJ/h
Q
          : heat load on the generator, kJ/day
          : heat abstracted from the biogas, kJ/m3
```

r : radius of circular fin equivalent to square

fin, m

 $r_i$  ,  $r_o$  : inside and outside radius of tube,  $m_i$ 

r\* : dimensionless parameter,  $(\frac{r + 3/2}{r_0})$ 

T : temperature, °C

T : maximum ambient temp. (= 41.3°C for Kanpur city)

 $\Delta T_{m}$ : LMTD, log mean temperature difference, °C

TR : tonnage of refrigeration

U : overall heat transfer coeff., unfinned,kJ/m2-h-°C

uf : overall heat transfer coeff., finned, kJ/m2-h-°C

V : volume of biogas, m<sup>3</sup>/day

x : concentration of LiBr in water, percent

: quality of water vapour

Non-dimensional Parameter

Gr : Grash of number,  $\frac{1^3 \rho^2 \beta g. T}{\mu^2}$  or  $\frac{d^3 \rho^2 \beta T}{\mu^2}$ 

Gz : Graezt number, Re Pr (d/L)

Pr : Prandtle number,  $C_{p} \mu/k$ 

Re : Reynolds number,  $d G/\mu$ 

Nu : Nusselt number,  $\frac{hd}{k}$  or  $\frac{h1}{k}$ 

### GREEK LETTERS

```
Fin thickness, m
δ
               Wall thickness (in the conduction of heat
8
               from inner to outer surface), m
              Parameter defined: 1 1/h/k(6/2)
Ψ
            : Fin effectiveness
               Stephan Boltzman constant = 20.42 \times 10^{-8} \frac{kJ}{m^2 - h - K^4}
 σ
                emmissivity,
            : 1.0 for a black surface
€~
               dynamic viscosity, kg/m-h
μ
            : density, kg/m<sup>3</sup>
ρ
            : temperature difference, °C
AT
              coefficient of thermal expansion, 1/T, (1/K)
В
               kinematic viscosity, (\mu/\rho), m^2/h
° C
              degree Celcius
K
              degree Kelvin
σt
            : surface tension, N/m
            : precooler and preheater effectivenesses.
\epsilon_1, \epsilon_2
```

respectively  $(\epsilon = \epsilon_1 = \epsilon_2)$ 

# SUBSCRIPTS

a : absorber side

c : condenser side

e : evaporator side

f : fin

g : generator side

go : optimum generator

i : inside

1 : liquid

o : outside

r : radiative

s : saturated

sp : superheat

v : vapour

w : wall

x : helix

#### ABSTRACT

A pumpless LiBr-H<sub>2</sub>O vapour absorption system has been designed to operate using biogas as a source of energy. The capacity of the system is selected on the basis of the requirement of an average agricultural family. Optimum generator temperatures for different operating conditions have been obtained based on economic criteria and are shown graphically. The design of the system components is done for the optimum generator temperature corresponding to the generator pressure of 80 mm of mercury and the evaporator temperature of 5°C. The conditioned space can be maintained at about 7 to 8°C. The heat rejection from the system is by natural convection of air. The temperature difference between the maximum ambient temperature and the designed condition for heat rejection to the former is about 5 to 6°C.

The present work would be most helpful in the design of a refrigerator, water cooler or an air-conditioner for different operating conditions. The system designed would serve the agrobased rural community in preserving eatables, vaccines, vital medicines, etc.

The effects of effectivenesses on biogas cost have been studied. The variation in biogas cost with system capacity has been also studied. Multipliers were computed for evaporator temperatures other than 5°C such that the same would enable the users to utilize the present results for a wide range of operation.

Components of the refrigeration system have also been fabricated and assembled. In the limited time inspite of best efforts, it was not possible to make the assembly completely leakage-free, which is essential for sustained running of the set-up and experimentation.

The detailed method for the design of various components would enable users to design the system components to suit their requirement.

#### CHAPTER - I

#### INTRODUCTION

#### 1.1 Social Rammifications

The attractive city life and poor village management is causing heavy influx of people from rural areas to cities. This is posing threat to appropriate urban life and is becoming totally beyond the manageable limit of various authorities. As such if the living standard and job opportunities (need of the day in the national interest) are improved in the rural areas, there will be tremendous reduction in the rural outflux to cities. The reduction in influx to city would also reduce traffic load and urban housing shortage.

#### 1.2 Energy Demand

Presently, the escalating cost of energy and impending fuel shortages have created interest in the research and development of alternate energy resources. A few such promising sources which have been under active investigation and development [1,2,3,4,5], are the solar energy, blomass energy, wind energy, tidal energy, waste heat recovery, etc.

#### 1.3 Rural Applications

A wide spectrum of work is being done in the field of solar energy. Thus solar powered cookers, heaters, refrigerators, air conditioners, etc. are gaining market day by day. The biomass energy is a good source specially for agrobased rural areas and at places where electricity failure is frequent. Our country is an agricultural land and the cultivation is mostly done using oxen, and for milk buffaloes and cows are tamed. It means the availability of animal dung is a continuous and everlasting source. If biogas is produced from the dung, it would provide better rural life. The fermentation . increases manure value of animal dung by as high as 40% [6]. This would cause considerable curtailment in synthetic fertilizer. Moreover the use of natural fertilizer retains better productivity of land. Also, the extraction of the biomass energy, which is mostly from the organic materials, being wasted, solve the problem of pollution. Use of biogas also reduces demand for wood etc. as cooking fuel, which saves the trees from being cut in large scale. Biogas has been widely used for cooking and lighting. The hybridization of biogas has been proposed in [7] to serve as a standby power source for the solar powered vapourabsorption refrigeration system. A family may need a 0.1 ton capacity refrigerator operating for about 12 hours

per day (at generator pressure,  $p_g = 80 \text{ mm of H}_g$ , evaporating temperature,  $T_e = 5^{\circ}\text{C}$  with COP = 0.746) using 1.32 m³/day of gobar gas. For a normal family, cooking gas required/day is 1.4 m³/day and for lighting is 1.2 m³/day. Thus, total gas required turns out to be approximately 4 m³/day. An average animal dung (wet) yield [8] is about 15 kg/day, giving about 0.04 m³ of biogas per kg of dung, requiring about 6 to 7 animals, which is normally within the scope of a good agricultural family.

#### 1.4 The Refrigeration System

The vapour-absorption system eliminates the tremendously large compression energy, owing to the reason that the system operates at low pressures, requiring negligible pumping energy. The special feature of the vapour-absorption is the use of thermal energy, a low grade energy as a byproduct of waste heat from the process industry, blast furnace, steel plants, automobile exhausts, solar energy or biogas energy. Further, its COP can be kept high even if the evaporator temperature falls. A comparative study [9] for a single stage refrigerant absorbent combination is given in table 1.1.

Table 1.1

Comparative study of various refrigerant-absorbent combinations [9]

Absorbent- refrigerant combination	NH 3	<sup>H</sup> 2 <sup>O</sup>	NH 3	-Lino 3	NH3-Nascn	Sink   Tempera-   ture   T_ = 30°C
Generator temperagure T <sub>g</sub> , °C	115	140	90	115	100	c
Evaporating temperature, °C	-10	<b>-</b> 25	-10	<b>-2</b> 5	-10	

Available fluid temperature from the flat plate collector [9] covers a range from approximately 80°-115°C, but are generally below 90°C. It can be used to run systems for which generation temperature is not large. Also waste heat available is at a temperature of about 100°C.

It has been reported [10,11,12] that LiBr-H<sub>2</sub>O is the best combination for the optimum performance of vapour-absorption system temperature above 4°C. The COP of this system varies from 0.6 to 0.75, whereas, the COP of NH<sub>3</sub>-NaSCN and NH<sub>3</sub>-H<sub>2</sub>O systems vary from 0.11 to 0.27 and 0.05 to 0.14, respectively.

#### 1.5 Economic Factors

The limitation in the availability of the temperature from the solar collectors necessitates the use of

other sources yielding higher temperatures. Biogas is one workable at a wide range of temperatures; limit being put to the available capacities of biogas plants. India biogas units of 2,3,4 and 6 m<sup>3</sup>/day have been widely used. Also, the cost comparison between the available energy resources and the renewable ones show that the initial investment is quite high, but in the long run it is comparable with other sources. It is reported in [8] that the energy from 1 m<sup>3</sup> of biogas is equivalent to that of 0.62 litre of kerosene oil and 4 kg of firewood. cost of 1 m<sup>3</sup> of biogas is estimated to be Re.O.72 per day if the life of biogas plant is taken as 15 years. In the same way the cost of biogas required in the refrigerating unit of 0.1 ton capacity (for 1.32 m3/day) is about Re.0.84 per day, whereas, cost of electrical power in the vapourcompression system for the same capacity and time of operation is about Re. 0.66 per day (i.e., 0.048 kW x 12 hour x Re. 0.55). For higher capacities the cost of biogas reduces considerably. This comparison, obviously does not have any meaning where electricity is rarely available or not available at all.

#### 1.6 Present Work

The objective of the present work is to develop a cooling unit which would preserve eatables at about 7 to 8 °C especially for the agrobased areas using biogas

as a source of energy. With this view economic cost analysis of the biogas required in the generator of the LiBr-H20 vapour-absorption system has been done at various operating conditions, and the optimum generator temperatures have been evaluated correspondly. The optimization has been carried out for NH2-H20 combination using flat plate collector as a source of energy [13]. LiBr-H,0 system using solar energy has been optimized with respect to maximum COP [7]. It reports for  $p_q = 71.9$  mm of  $H_q$  and  $T_e = 5$ °C, optimum generator temperature,  $T_{g_0} = 93$ °C (using precooler and preheater both,  $\epsilon = 0.8$ ) and  $T_g = 103.5$ °C (for  $\varepsilon = 0$ ). In the present work of optimization, the generator temperature  $T_q$  is varied keeping the generator pressure  $\mathbf{p}_{\mathbf{q}}$  , the evaporator temperature  $\mathbf{T}_{\mathbf{e}}$  and precooler and preheater effectivenesses as parameters. The COP and the available energy to the generator,  $\Omega_{\text{gas}}$  are functionally related to  $T_q$ . The yearly running cost and the capital cost for biogas plant are functionally related to T using least square regression technique. Results are obtained for different refrigeration capacities as well.

The components of the refrigerating system have been designed on the basis of optimum values obtained at generator pressure 80 mm of  ${\rm H_g}$ ,  ${\rm T_e}$  = 5°C and  ${\rm ^6_2}$  = 0.75 (only preheater is used). In the fabrication and assembly

of such a system having large number components, even slightest leakage hampers the experiment because of the vacuum (1 atm) to be maintained inside the system tubings. To realize a completely leakage free set-up, it normally requires considerable amount of time. The system has been fabricated and instrumented, but very minute leakages still remain to be rectified in order to obtain reliable experimental data.

#### CHAPTER - II

#### MATHEMATICAL FORMULATION

#### 2.1 System Selection

Since the aim is to develop a cabinet refrigerator or a cooling unit required in primary health care centres mainly for the agrobased areas, it is necessary that the system be simple, cheap and independent of electricity. The most common absorption refrigeration systems are NH<sub>3</sub>-H<sub>2</sub>O, LiBr-H<sub>2</sub>O, NH<sub>3</sub>-NaSCN and NH<sub>3</sub>-LiNO<sub>3</sub>. Comparative study among these show [10,11,12] that LiBr-H<sub>2</sub>O absorption system would be the best amongst all. In this system water acts as a refrigerant and Lithium browide (salt) as an absorbent. Few attractive features of LiBr-H<sub>2</sub>O system are presented below:

- a) Higher COP values, about 0.6 to 0.75.
- b) Pure water vapour leaves the generator since LiBr does not evaporate. Hence an analyzer and a rectifier are eliminated giving a simple system.
- c) Purging of water from the evaporator is trivially eliminated.

- d) Overall working pressure in the system is barely above 0.15 bar (100 mm of  $H_g$  absolute), there is no need to design the system along high pressure codes.
- e) The pressure inside the system is below atmosphere, hence no fear of bursting or accidental hazard.
- f) Water being non-toxic and nonflammable, the system can be used directly for chilling coil for the air conditioners.
- ween the difference in the operating pressures between the generator and the evaporator is very small, the pumping energy required is negligible. Moreover, pumping mechanism is eliminated simply by the adjustment of the elevation by a metre or so of the absorber, generator and the evaporator; whereas, the elevation has to be increased by 100 metres to achieve the operating pressure difference in NH<sub>3</sub>-H<sub>2</sub>O system. Thus, LiBr-water system is free from vibration and noise as well. Hence no strong foundation required.
- h) No leakage in the system, since shaft seal is absent.
- i) Cost of the system is very small since there are no moving parts, glands, valves, etc.
- j) No electrical energy required.
- k) Water having higher enthalpy of vaporization than

NH<sub>3</sub> is used. The refrigerant is almost free of cost. With these advantages the LiBr-H<sub>2</sub>O system is selected for the present work.

#### 2.2 System Description

Absorption cycles are heat operated in which a secondary fluid, the absorbent, absorbs the primary fluid, gaseous refrigerant from the evaporator maintained at low pressure. Absorption takes place because of more affinity of absorbent for the refrigerant. To augment the absorption in small volume of absorber, the large contact area is provided. The energy released [14] during the absorption process is due to heat of condensation, sensible heat and heat of dilution, which is rejected to a sink.

The refrigerant-absorbent solution is pumped into the generator via a heat exchanger where the refrigerant and the absorbent are separated by a distillation process, because at higher temperature the absorbent has less affinity for the refrigerant molecules. If the pure absorbent is non-volatile, as in LiBr-H<sub>2</sub>O system, a simple still is adequate. But in systems using volatile absorbents, like water in NH<sub>3</sub>-H<sub>2</sub>O, would require additional distillation equipments such as analyzer and rectifier. If the refrigerant is not essentially free of absorbent,

vaporization in the evaporator is hampered. If the absorbent material tends to become solid, as in LiBr-H<sub>2</sub>O system, enough refrigerant must be present to keep the pure absorbent material in the dissolved form at all times; and can be checked by avoiding excessive heating in the generator.

Figure 2.1(a) exhibits the vapour-absorption system having different components. The cycle with state points is shown in Fig. 2.1(b). The weak solution of LiBr-H<sub>2</sub>O from absorber is pumped into the generator through a preheater (PH) in order to reduce heat transfer,  $\Omega_{\rm g}$ , to the generator. The strong solution returns to A through the preheater and throttle valve. The refrigerant vapour from the generator is condensed in C and then throttled to the evaporator E. The condensate gets subcooled in the precooler (PC) by the evaporator vapour. Thus, the refrigeration effect gets enhanced. The refrigeration effect  $\Omega_{\rm e}$  is produced in the evaporator due to vaporization of water.

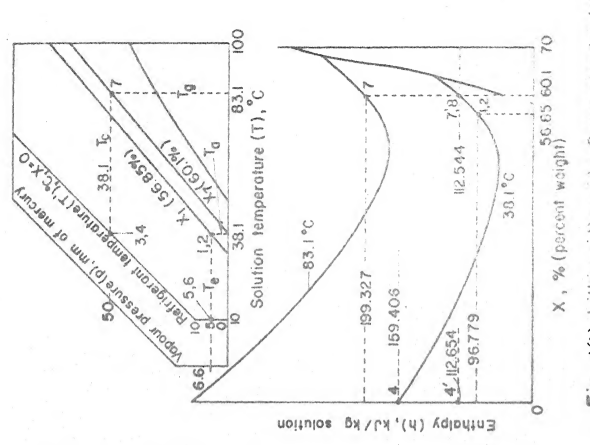
### 2.3 Mathematical Modelling

Absorber:

# 2.3.1 Thermodynamic analysis of the system

Referring to Fig. 2.1(a), the mass conservation in terms of concentration of the absorbent across absorber is given by:

$$m_6 \times_6 + m_7 \times_7 = m_1 \times_1$$
 (2.1)



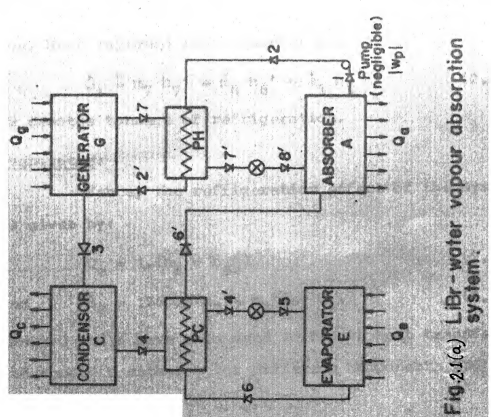


Fig. 2.1(b) Lithium Brounds Concentration (X)

where m and x denote for mass and concentration, respectively at different state points.

Since the pure vapour flows out of the evaporator,  $x_6 = 0$ . With  $m_6 = 1$  and  $m_7 = m_1 - 1$ . One gets from Eq. (2.1) as:

$$m_1 = x_7/(x_7 - x_1) \tag{2.2}$$

and 
$$m_7 = x_1/(x_7 - x_1)$$
 (2.3)

Also,

$$m_1 = m_6 \cdot m_1$$
 (2.4)

$$\mathring{\mathbf{m}}_{7} = \mathring{\mathbf{m}}_{6} \cdot \mathbf{m}_{7} \tag{2.5}$$

where m<sub>6</sub> is defined by Eq. (2.8).

Now, heat rejected from absorber per TR is:

$$\dot{Q}_{3} = \dot{m}_{7} h_{7}' + \dot{m}_{6} h_{6}' - \dot{m}_{1} h_{1}$$
 (2.6)

TR denotes tonnage of refrigeration.

# Evaporator:

Hence, the refrigeration effect of the system is given by:

$$Q_{e} = 1 \cdot (h_{6} - h_{4}') \tag{2.7}$$

and 
$$\dot{m}_6 = 12600/(h_6 - h_4')$$
 (2.8)

where Q and h have been used to denote heat transfer and specific enthalpy for different components and state points, respectively.

#### Generator:

Energy balance for the generator gives:

$$Q_{q} = m_{3} h_{3} + m_{7} h_{7} - m_{2}' h_{2}'$$
 (2.9)

Using: 
$$m_3 = m_4 = m_4' = m_5 = m_6 = m_6' = 1 \text{ kg}$$
  
 $m_1 = m_2 = m_2' \text{ ; } m_7 = m_7' = m_8$ 

One gets:

$$Q_q = [h_3 + (x_1 h_7 - x_7 h_2')/(x_7 - x_1)] (2.10)$$

Heat load in the generator per TR is:

$$\dot{Q}_{q} = \dot{m}_{3} \cdot Q_{q} \qquad (2.11)$$

where  $\dot{m}_3 = \dot{m}_6$ 

# Precooler:

The enthalpy of subcooled condensate, h4' is:

$$h_4' = h_4 - \epsilon_1 (h_6' - h_6)$$
 (2.12)

where  $\epsilon_1$  is effectiveness of the precooler.

# Preheater:

Similarly, for the preheater (effectiveness,  $\epsilon_2$ ):

$$h_2' = h_1 + \epsilon_2 (x_1/x_7) (h_7 - h_7')$$
 (2.13)

Hence,

$$COP = Q_e/Q_g = (h_6 - h_4')/[h_3 + (x_1 h_7 - x_7 h_2')]/(x_7 - x_1)]...(2.14)$$

#### Condenser:

Heat rejection from the condenser per TR is:

$$\dot{\Omega}_{C} = \dot{m}_{3} (h_{3} - h_{4})$$
 (2.15)

The thermodynamic properties of LiBr-H<sub>2</sub>O solution of [15] have some inconsistency. The proper units have been taken from [14], which are in terms of operating temperature, pressure and concentration of the solution, given in Appendix A. Also, the properties of pure water [16] are listed in A. The calculations are made in S.I. units.

# 2.3.2 Functional Relation for COP [13]

Using appropriate equations and properties,  $\operatorname{COP}$ 's were computed at different generator temperatures for a set of operating parameters such as the generator pressure being equal to that of condenser, the evaporator temperature and precooler and preheater effectiveness. Thereafter  $\operatorname{COP}$ 's were functionally related to  $\operatorname{T}_g$  using least square error regression method for each set.

i.e., 
$$COP = \sum_{j=0}^{j=9} a_j T_g^j = F_1(T)$$
 (2.16)

# 2.3.3 Useful Energy from Biogas

Composition of the biogas was taken as [3,6]: (Methane)  $\text{CH}_4$  - 55% and (Carbondioxide)  $\text{CO}_2$  - 45% ... (2.17)

Combustion equation can be written as:

0.55 
$$\text{CH}_4 + 0.45 \text{CO}_2 + \text{a} (\text{O}_2 + 3.76 \text{N}_2) \longrightarrow$$
 $\text{CO}_2 + 1.1 \text{H}_2\text{O} + \text{bO}_2 + 3.76 \text{a} \text{N}_2$  (2.18)

where a and b are constants.

In order to promote better air fuel mixing 10% excess air is used. Hence,

$$\frac{b}{1+b+3.76 \text{ a}} = 0.1 \text{ or } 0.9 \text{ b} = 0.1 + 0.376 \text{ a}$$

$$(2.19)$$

$$0_2$$
 balance gives:  $0.45 + a = 1 + 0.55 + b$  .... (2.20

Solving Eqs. (2.19 and 2.20), we have, a = 2.08015 and b = 0.98015.

Air fuel ratio,

A/F = 
$$\frac{\text{mass of (2.08015 mole of O}_2 + 7.82136 \text{ mole of N}_2)}{\text{mass of (0.55 mole of CH}_4 + 0.45 \text{ mole of CO}_2)}$$
.... (2.21)

1 mole of  $\rm O_2$  ,  $\rm N_2$  ,  $\rm CH_4$  and  $\rm CO_2$  equal to 32, 28, 16 and 44 respectively. Substituting respective values in Eq. (2.21), we have

$$A/F = 9.9847$$

For a combustion reaction occurring at constant pressure in steady flow, the first law of thermodynamics with enthalpy of combustion can be written as [17],

$$Q_{gas} = \Delta H_s - [Q_{prod.} - Q_{react.}]$$
 (2.22)

$$Q_{gas} = \Delta H_{s} - \left[ \sum_{p} n_{p} \left( \overline{h}_{p} \left( T_{p} \right) - \overline{h}_{p} \left( T_{sd} \right) \right] - \sum_{r} n_{r} \left( \overline{h}_{p} \left( T_{r} \right) - \overline{h}_{p} \left( T_{sd} \right) \right) \right]$$

$$(2.23)$$

where,

 $\Delta H_{s}$  = Enthalpy of combustion.

gas = Heat abstracted from the fuel during combustion
process.

 $\bar{h}_p$  ,  $\bar{h}_r$  = Specific enthalpies of products and reactants, respectively (kJ/kg- mole)

 $T_{p}$  ,  $T_{r}$  = Temperature.

T<sub>sd</sub> = Standard temperature, usually 25°C (298.15 K)

Since, only CH $_4$  burns in the biogas, enthalpy of combustion, from table A-7 [17] at T $_{\rm sd}$  = 298.15 K,  $\Delta$ H $_{\rm s}$  = - 11,946 x 4.187 kJ/kg. For 55% methane as reactant,

$$\Delta H_{s} = 4,40,158.375 \text{ kJ/kg-mole}$$

Rewriting Eq. (2.23) for the combustion equation, Eq. (2.18), we have

$$\begin{split} Q_{\text{gas}} &= \Delta H_{\text{s}} - \left[ \left( \left\{ \bar{h}_{\text{CO}_{2}} \left( T_{\text{p}} \right) - \bar{h}_{\text{CO}_{2}} \left( T_{\text{sd}} \right) \right\} \right. \\ &+ 1.1 \times \left\{ \bar{h}_{\text{H}_{2}\text{O}} \left( T_{\text{p}} \right) - \bar{h}_{\text{H}_{2}\text{O}} \left( T_{\text{sd}} \right) \right\} \\ &+ 0.98015 \times \left\{ \bar{h}_{\text{O}_{2}} \left( T_{\text{p}} \right) - \bar{h}_{\text{O}_{2}} \left( T_{\text{sd}} \right) \right\} \\ &+ 7.82136 \times \left\{ \bar{h}_{\text{N}_{2}} \left( T_{\text{p}} \right) - \bar{h}_{\text{N}_{2}} \left( T_{\text{sd}} \right) \right\} \right] \end{split}$$
 Eq. contd....

$$- (0.55 \times \{\bar{h}_{CH_{4}}^{(T_{r})} - \bar{h}_{CH_{4}}^{(T_{sd})}\}$$

$$+ 0.45 \times \{\bar{h}_{CO_{2}}^{(T_{r})} - \bar{h}_{CO_{2}}^{(T_{sd})}\}$$

$$+ 2.08015 \times \{\bar{h}_{O_{2}}^{(T_{r})} - \bar{h}_{O_{2}}^{(T_{sd})}\}$$

$$+ 7.82136 \times \{\bar{h}_{N_{2}}^{(T_{r})} - \bar{h}_{N_{2}}^{(T_{sd})}\}$$

$$\cdots (2.24)$$

Let us assume that the reactants burn at room temperature (i.e.,  $T_{am} = T_r = 314.45 \text{ K}$ ) and the products Leave at 50°C above the generator temperature. Let us assume that the generator is operating at a temperature,  $T_g = 116$ °C, so that

$$T_p = T_q + 50 = 439.15 \text{ K}$$

Hence, the specific enthalpies of the various compounds at the corresponding temperatures  $T_p$ ,  $T_r$  and  $T_{sd}$ , listed in table 2.1, are taken from table A-9 [17].

Table 2.1 Specific Enthalpies kJ/kg-mole of Various Compounds at Typical Value of  $T_p$  ,  $T_r$  ,  $T_{sd}$ 

Temperature(K)	T	Ť	<u> </u>	
Compound	439.15	314.45	<sup>1</sup> sd 298 <b>.1</b> 5	
co <sub>2</sub>	15,059.13	9,934.81	9,379.34	
H <sub>2</sub> O	14,727.68	_	8,471.37	
02	2,897.05	9,097.61	8,668.67	
N <sub>2</sub>	12,813.47	9,102.56	8,678.69	
CH <sub>4</sub>		8,451.46	8,050.49	

Substituting the respective values of specific enthalpies from table 2.1 in Eq. (2.24) we have

$$Q_{gas} = 3,95,790.56 \text{ kJ/kg-mole}$$

Using equation of state at normal temperature and pressure

$$p\overline{V} = \overline{R}T \tag{2.25}$$

with, p =  $1.013 \times 10^5 \text{ N/m}^2$ , T = 314.45 K and  $\overline{R}$  = 8.317 kJ/kg mole K

$$\overline{V} = 25.7 \text{ m}^3/\text{kg-mole}$$

Making use of the specific volume  $\overline{V}$ ,

$$Q_{gas} = 15,400.41 \text{ kJ/m}^3$$

Similarly, values of  $Q_{gas}$  were calculated for different operating temperatures and were functionally related to  $T (= T_{g} + 50)$  as:

$$\Omega_{\text{gas}} = 17.560.37152 - 12.64482465 T$$

- 
$$(1.7546455 \times 10^{-3}) \text{ T}^2 = \text{F}_2(\text{T})$$
 (2.26)

# 2.3.4 Biogas Cost with Volume

From [8], the capital cost of biogas plants and their annual expenditure along with the annual income as reported in "An Economic Analysis of a 3m<sup>3</sup> Gas Plant conducted by the Khadi and Village Industries Commission in India (1978)" [18] is estimated and is functionally related to the biogas volume using the least square regression

method. The capital cost includes the total investment on construction of the biogas plant, piping, stove, etc. The interest on the investment is taken as 9% per annum, depriciation on structure, piping and stove as 5% per annum and cost of painting each year is taken as Rs. 50/litre, constitute the annual expenditure. The annual income includes the cost of gas (taken as Rs. 185/m³) and the selling price of manure is Rs. 44/ton. The cost of gobar is taken as 1.5 times the selling price of manure, which also adds to the annual expenditure. It is reported [8] that 1 m³ biogas plant requires 25 kg of gobar, and out of 23 ton gobar, 7 ton is composted and 16 ton is refused. (i.e., useful manure). The effective working days for the biogas plant are assumed as 325 days/year. The cost-volume relation is represented as follows:

$$C_1 = B_1 V^{n_1}$$
 (2.27)

and

$$C_2 = B_2 v^{n_2}$$
 (2.28)

where,

$$B_1 = 1597.48672, n_1 = 0.706043244$$
 (2.29)

$$B_2 = 220.1583947$$
,  $n_2 = 0.546585428$  (2.30)

From Eqs. (2.7, 2.8 and 2.14), the heat load on the generator can also be written as:

$$Q_{q}' = TR \times 12,600 \times 24 / COP$$
 (2.31)

Hence, volume of the biogas required for a given capacity and COP of a refrigerating system can be found as:

$$V = Q_{g}'/Q_{gas} = TR \times 12600 \times 24 / (COP \times Q_{gas})$$

$$(2.32)$$

Assuming an operating condition of 70%, we have from Eqs. (2.16, 2.26 and 2.32),

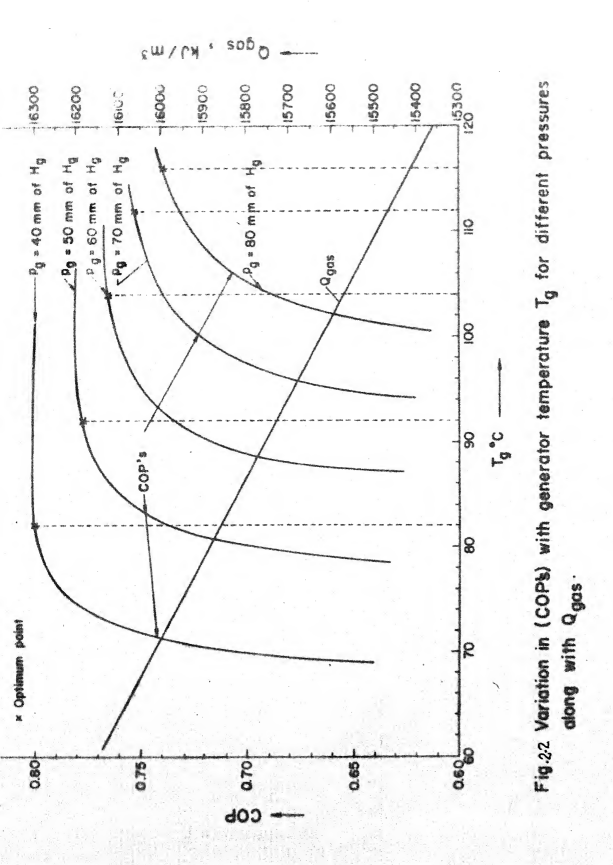
$$V = TR \times 12600 \times 24 \times 0.7 / (F_1(T) \times F_2(T))$$

$$= 211680 \times TR / F(T) \qquad (2.33)$$
where, 
$$F(T) = F_1(T) \times F_2(T) \qquad (2.34)$$

$$2.4 \quad \text{Optimization}$$

The total cost of the whole system is the objective function related to generator temperature for a set of parameters. The generator temperature should be such that the cost of biogas energy required to operate it is minimum. From Eqs. (2.27 and 2.28) cost of biogas is directly proportional to its volume. Hence, for minimum cost the minimum volume of biogas should be used. From Eq. (2.34) volume of biogas required is minimum if F(T) is maximum, that is, the product of COP and  $Q_{\rm gas}$  is maximum. The variation in COP of the system for different pressures along with  $Q_{\rm gas}$  with generator temperature is shown in Fig. (2.2).

If  $C_2$  exists at the end of each year of L years and R is the interest rate, the present worth, P is obtained



by summing the present worth of each of the payments of amount  $\mathbf{C}_2$  .

Thus, 
$$P = \frac{C_2}{L} \sum_{i=1}^{L} (1/(1+R)^{L-1})$$
 (2.35)

Eq. (2.35) being geometric progression of constant ratio 1/(1+R),

then,

$$P = \frac{C_2}{L} \left[ ((1 + R)^L - 1)/(R(1 + R)^{L-1}) \right]$$
 (2.36)

The total cost of biogas per year is given by

$$C_{T} = \{C_{1} + C_{2} [((1+R)^{L} - 1)/(R(1+R)^{L-1})]\} /L$$

$$(2.37)$$

The optimum value yielding  $C_{\underline{T}}$  minimum was computed using iterative method. The computer results for the same are given in Appendix-B.

### CHAPTER - III

#### LOAD CALCULATIONS

### 3.1 Cooling Load Calculations

Perishable commodities for usage in day to day life are sometimes preserved for convenience to avoid spoilage. To produce comfort condition during hot season days refrigeration is required. For these purposes LiBr-H<sub>2</sub>O system has been selected. The capacity of the system is decided on the basis of cooling load calculations. The details are as follows:

- (i) Cooling load due to commodities
  - (a) continuous commodity load
  - (b) intermittent commodity load.

# (ii) Structure load.

Cooling load due to commodities:

The heat release from the commodity comprises sensible cooling of products from the ambient temperature to freezing temperature, enthalpy of freezing, sensible cooling below freezing and heat of respiration [19]. Since water is used as a refrigerant, the system would operate above freezing temperature. Hence, the heat

release from the commodities for storage above 0°C reduces to,

$$Q_{cod} = M \left[ C_{p} \left( T_{am} - T_{store} \right) \right] / \left( CF \times t \right) + M \cdot hr/1000$$

$$\dots \qquad (3.1)$$

where

 $Q_{cod}$  = heat release from the commodity in kJ/h

M = amount of the commodity stored in kg

c
p = specific heat of the commodity above
freezing

T<sub>am</sub> = ambient temperature (314.45 K)

T<sub>store</sub> = storage temperature (278.15 K)

CF = chilling rate factor

t = chilling time

hr = energy release due to respiration per tonne mass per unit time.

The continuous commodity loads are due to the eatables which are stored for all 24 hours, such as, apple, tomato, crated eggs, fish, carrot, oranges, etc. The intermittent loads are due to commodities which kept in the refrigerator at times, such as, milk, curd, water, etc. The parameters in Eq. (3.1) for different commodities are tabulated in table 3.1. The cooling load due to different commodities,  $\Omega_{\rm cod}$ , is also tabulated. The values of parameters in table 3.1 are taken from [19,20,21].

Table 3.1

Design data for different commodities along with cooling load required for their storage

Parameters Commodities	M (kg)	C p (kJ/kg°C)	CF	t. (h)	hr (kJ/ton-h)	Ω cod (kJ/h)
Apple	3	3.768	0.67	24	26.166	25.0
Tomato	1	3.85	1.0	24	54.26	5.88
Eggs(crated)	0.65	3.192	0.85	10	155.02	8.89
Fish	2	3.182	1.0	4	245.448	58.244
Carrot	2	3.6	0.8	24	122.103	13.857
Oranges	. 3	3.768	0.7	22	53.303	26.805
Milk	5	3.906	0.85	5	86.122	167.240
Curd	4	3.906	0.9	6	86.122	106.24
Water	5 h	am=171.75	1.0	3	O	502.566
	h	= 20.98 store	3			

where,  $h_{am}$  and  $h_{store}$  are the enthalpies of water corresponding to temperatures  $T_{am}$  and  $T_{store}$ .

The total cooling load on summing the individual commodity loads comes out to be:

$$\bar{Q}_{cod} = 914.72 = 915 \text{ kJ/h}$$

Structure load:

The size of the storage cabinet is assumed to be same as that of a 165 litre refrigerator.

Dimensions of the refrigerator:

Outer : 0.55 x 0.55 x 0.92 m

Inner :  $0.45 \times 0.45 \times 0.82 \text{ m}$ 

Insulation: 0.05 m (glass wool).

Overall heat transfer coefficient for the cabinet is given by,

$$U_{ob} = \frac{1}{h_{i_b} + \frac{1}{h_{o_b}} + \frac{\delta' woo1}{k_{wool}}}$$
 (3.2)

where  $h_{ib}$  and  $h_{ob}$  are respectively inner and outer heat transfer coefficients and are taken as:  $h_{ib} = h_{ob} = 100$  kJ/hm<sup>2</sup>°C and  $k_{wool}$  = thermal conductivity of the wool (= 0.15 kJ/hm<sup>2</sup>°C).  $S_{wool}$  = insulation thickness Substituting respective values in Eq. (3.2)we have

$$U_{O_h} = 2.83 \text{ kJ/hm}^2 \circ \text{C}$$

Also

$$Q_{s} = U_{o_{b}} A_{i_{b}} (T_{am} - T_{store})$$
 (3.3)

where,  $Q_s$  is the structure load and  $A_i$  is the heat transfer area ( $\simeq$  4 x 0.45 x 0.82 + 2 x 0.45 x 0.45).

Substituting the respective values in Eq. (3.3), we have

$$Q_{s} = 193.233 \text{ kJ/h}$$

Increasing by 10% for safety,  $Q_s = 213 \text{ kJ/h}$  Hence, total cooling load =  $\overline{Q}_{cod} + Q_s = 1128 \text{ kJ/h}$  This load is the peak load, i.e. if all the commodities listed in table 3.1 are kept in the cabinet at a time. But the actual load will be fluctuating according to the requirement as shown in Fig.(3.1). From Fig. (3.1)  $Q_{area} = 476 \text{ kJ/h}$ , but the maximum load as can be seen from the Fig.(3.1) is at 10 to 13 hours; on the basis of which the cabinet is to be designed,

i.e.,  $\dot{\Omega}_{\rm e}$  = 593.333 kJ/h  $\simeq$  600 kJ/h where  $\dot{\Omega}_{\rm e}$  is the rate of cooling load on the cabinet.

## 3.2 Energy Exchange for Components

Making use of the LiBr - concentration charts and equations the various parameters at pressure,  $p_g = p_C$  = 80 mm of  $H_g$  and evaporator temperature,  $T_e = 5^{\circ}C$  ( $p_e = p_a = 6.6$  mm of  $H_g$ ) are calculated and listed below: Table 3.2

Property values at optimum generator temperature  $(T_g)$  corresponding to  $p_g$  = 80 mm of  $H_g$ ,  $T_e$  = 5°C and  $\epsilon$  = 0.75

Tg_	T <sub>c</sub> =T <sub>a</sub>	× <sub>1</sub>	× <sub>7</sub>	h <sub>1</sub> =h <sub>2</sub>	h <sub>3</sub>	h <sub>4</sub>	h <sub>6</sub>	h <sub>7</sub>	h <sub>7</sub>
° C	°C		*,0					kJ/kg	

116 47 61.4 69.0 136.8 2717.4 196.9 2510.1 300.1 182.5

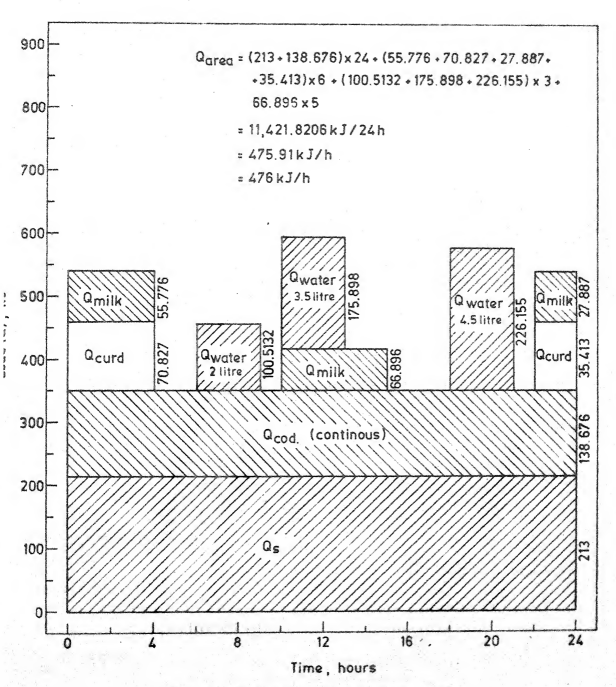


Fig. 3.1 Daily variation of cooling load.

The energy exchange due to different components, using Eqs. (2.1 to 2.15) and substituting the respective values from table 3.2, are as follows:

The capacity of the refrigerating unit, as calculated above is  $\dot{\Omega}_{\rm e}$  = 600 kJ/h

## Evaporator:

From Eq.(2.7) we have,

 $Q_e = 2313.156$  kJ/kg and hence,  $\dot{m}_6 = \dot{Q}_e/Q_e = 0.26$  kg/h Absorber:

From Eqs. (2.4 to 2.6) we have,

 $\dot{m}_1 = 2.361 \text{ kg/h}, \dot{m}_7 = 2.101 \text{ kg/h} \text{ and } \dot{Q}_a = 715 \text{ kJ/h}$ 

Generator:

From Eqs. (2.10 and 2.11) we have,

$$\mathring{m}_3 = \mathring{m}_6$$
 and  $\mathring{Q}_g = 830 \text{ kJ/h}$  From Eq. (2.14),  $\text{COP} = \frac{Q_e}{Q_g} = \frac{\mathring{Q}_e}{Q_g} = 0.72$  TR =  $\frac{\mathring{Q}_e}{12600} = 0.048$ 

Condenser:

Eq. (2.15) gives,

 $\dot{Q}_{c} = 656 \text{ kJ/h}$ 

3.3 Biogas Requirement for the System

From Eq. (2.26), the useful gas required at the

optimum generator temperature,  $T_{g_0} = 116$ °C is given as:  $\Omega_{gas} = 15,400.41 \text{ kJ/m}^3$ 

Hence, the volume of gas is given by Eqs. (2.33 and 2.34) as:

 $V = 0.048 \times 211680 / (0.72 \times 15.400.41) = 0.92 \text{ m}^3/\text{day}$ 

#### CHAPTER - IV

#### SYSTEM DESIGN

The design of the vapour-absorption refrigeration system components require considerable informations such as feasibility of certain systems physical realizability, economic worthwhileness, financial feasibility, operating environmental conditions including chemical and physical reactions, etc. Besides these, one has to consider factors such as materials, components and their optimum combination and adaptability by customers. The cooling and component load calculations have already been discussed in Chapter-3.

### 4.1 Condenser Design

The condenser is the heat rejecting unit in a refrigerating system. When the saturated vapour comes in contact with a surface, at a temperature lower than the former, condensation occurs either in drops or as a film, [14]. The rate of heat flow depends on the condensate film thickness, which depends on the rate at which vapour is condensed and the rate of condensate removal. In the present case air cooled condenser has been selected due to simplicity. The condensate covers the inside tube

surface, flows by gravity, collects at the bottom of the inside horizontal tubes and flows by hydrostatic gradient. Chato [22] has obtained the following expression for condensation of refrigerants at low vapour velocities inside horizontal tubes as:

$$h_{i_{c}} = 0.555 \left[ \frac{\rho_{1_{c}}(\rho_{1_{c}} - \rho_{v_{c}}) g k_{1_{c}}^{3} h_{1v_{c}}}{\mu_{1_{c}} d_{i_{c}} (T_{s_{c}} - T_{w_{i_{c}}})} \right]^{0.25}$$

$$\dots (4.1)$$

in the range of

$$Re_{C} = \frac{d_{i_{C}}^{G_{V_{C}}}}{\mu_{C}} < 35,000$$

where Re is evaluated at inlet conditions to the tube and

$$h'_{lv_c} = h_{lv_c} + 0.68 C_{p_{w_c}} (T_{s_c} - T_{w_{i_c}})$$
 (4.2)

where

 $h_{lv_c}$  = latent heat of condensation

The tube and fin dimensions, selected, are tabulated in table 4.1.

		Table 4.1			
Tube	and Fin	dimension	o£	the	condenser

No. of Land Conference Survey Supplement	T	ube di	mensio	ns	Ŏ Ŏ	Fin d	imensions
(m)	(m)		(m)	(m)	ī <sub>c</sub> (m) x10 <sup>-2</sup>	•	6 <sub>C</sub> (m) <sub>2</sub> x10 <sup>-3</sup>
1.43	1.67	0.715	0.835	0.12	2.8675	4.17	5 0.42

The heat-transfer coefficient for the water-vapour flow on the inside of the tube is determined from the flow conditions with properties evaluated at bulk temperature. The free-convection heat-transfer coefficient on the outside of the tube depends on the temperature difference between the surface and the ambient air. The heat transfer area is increased by using fins.

The properties of saturated water at  $T_{s_c} = 47^{\circ}C$  [23,24] are given by table 4.2.

Table 4.2

Description of the second of t				4779.07	
Fropercie	es or sav	curated w	vater at T <sub>s</sub> c	= 4/°C	
k <sub>l</sub> c	μ <sub>lc</sub>	μ <sub>V</sub> c	Pla Pva	h <sub>iv<sub>c</sub></sub>	C <sub>pwc</sub>
(kJ/m-h-°C	)(kg/m-h	)(kg/m-h	1)(kg/m <sup>3</sup> )(kg/m	<sup>3</sup> )(kJ/kg)	)(kJ/kg°C)
2.322	2.232	0.0468 (at 116°C)	989.61 0.071	2391.2	4.1868

The Reynold number is

$$Re_{a} = \frac{d_{i_{C}} G_{V_{C}}}{\mu_{V_{C}}} = \frac{d_{i_{C}} (\mathring{m}_{3} / \frac{\pi}{4} d_{i_{C}}^{2})}{\mu_{V_{C}}} = 1179.574$$

where,  $\hat{m}_3$  = mass flow rate of water vapour in the condenser ( $\simeq 0.26 \text{ kg/h}$ )

and since Re < 35,000, using Eq. (4.1) and substituting values from tables (4.1 and 4.2) we have,

$$h_{i_c} = 8250.394 \left( h'_{1v_c} / (47 - T_{w_{i_c}})^{0.25} \right)$$
 (4.3)

and

$$h'_{1V_C} = 2391.2 + 0.68 \times 4.1868 (47 - T_{W_{i_C}}) (4.4)$$

Now, the temperatures,  $T_{w_i}$ ,  $T_{w_o}$  and  $T_f$  are unknown, hence iterative procedure is followed to find the former and hence, heat transfer coefficients  $h_i$ ,  $h_o$  and  $h_f$  are calculated.

Iteratively,  $T_{wic} = 46.97$ °C Hence, Eqs. (4.3 and 4.4) give:

••• 
$$h_{1v_c}' = 2391.2854 \text{ kJ/kg}$$
 and  $h_{1c} = 138.628.406 \text{ kJ/m}^2-h-\circ C$ 

Heat transfer coefficient on fin side having free convection is expressed as [25]

$$Nu = c (Gr \cdot Pr)^{n}$$
 (4.5)

The product Gr. Pr depend upon the fluid properties, the temperature difference between the surface and the fluid

and the characteristic length of the surface. The constant c and exponent n depend upon physical configuration and nature of flow.

Simplified equations for air as heat transport fluid, reported by [14, 23, 26, 27] are given below: For small cylinders, laminar range  $(10^4 < Gr \ Pr < 10^9)$ 

$$h_{oc} = 4.752 \left( \Delta T_{c} / d_{oc} \right)^{0.25}$$
 (4.6)

and for vertical plates (for finned surfaces), laminar range (same range as above)

$$h_{f_c} = 5.112 \left(\Delta T_c'' / \bar{l}_c\right)^{0.25}$$
 (4.7)

where, ho and hf are heat transfer coefficients outside the tube and the fin, respectively and  $\Delta T' = T_{W_0} - T_{am}$ ;  $\Delta T''_C = T_{f_C} - T_{am}$ 

Design ambient temperature for Kanpur city is  $T_{am} = 41.3^{\circ}C$ By iterative procedure I got  $T_{wo} = 46.96^{\circ}C$  and  $T_{fc} = 46.5^{\circ}C$ 

••• 
$$h_{c} = 20.3893 \text{ kJ/m-h-°C}$$
 and  $h_{fc} = 18.7593 \text{ kJ/m-h-°C}$ 

The overall heat transfer coefficient [28] is given by:

$$\frac{1}{U_{f_{O_{C}}}} = FF_{C} \left[ \frac{1}{h_{i_{C}}} + \frac{\delta c}{k_{W_{C}}} \right] + \frac{1}{h_{red_{C}}}$$

$$(4.8)$$

where,

and

$$A_{f_{C}} = 2 \pi n_{C} \left[ (r_{C} + c/2)^{2} - r_{O_{C}}^{2} \right]$$

$$A_{O_{C}} = 2 \pi L_{C} (1 - n_{C} c) r_{O_{C}}$$

$$A_{i_{C}} = 2 \pi r_{i_{C}} L_{C}$$

$$A_{i_{C}} = 2 \pi r_{i_{C}} L_{C}$$

$$h_{red} = (h_{f_{C}} p_{C} A_{f_{C}} + h_{O_{C}} A_{O_{C}}) / (A_{f_{C}} + A_{O_{C}})$$

$$\dots (4.10)$$

and thermal conductivity,

$$k_{W_C} = 1381.3 \text{ kJ/h-m-}^{\circ}\text{C}$$
 for copper tube wall.

Assuming, number of fins  $n_C$  mounted per unit length of the tube to be 72, the areas  $A_{i_C}$ ,  $A_{o_C}$  and  $A_{f_C}$  per unit length are calculated as:

$$A_{f_C} = 0.76495 \text{ m}^2$$
 ,  $A_{o_C} = 0.050878 \text{ m}^2$  ,  $A_{i_C} = 0.04492477 \text{ m}^2$  and  $FF_C = 18.16$ .

To account for the variation in the heat flux due to temperature difference at the tip and the root of the fin, fin effectiveness,  $\emptyset_{\rm C}$ , is defined. For constant thickness square fins the efficiency of a constant thickness annular fin of the same area can be used [29]. More accuracy, particularly with rectangular fins of large aspect ratio, can be obtained by dividing the fin into circular section [30]. From the fin efficiency curves

from Fig. (2.11)in [28],  $\emptyset_{\rm C}$  are found for the particular values of  ${\bf r}_{\rm C}^{*}$  and  $\Psi_{\rm C}$ 

where,

$$r_{C}^{*} = \frac{r_{C} + 6 c^{2}}{r_{O}}$$
 (4.11)

$$\Psi_{C} = \overline{I}_{C} \sqrt{h_{f_{C}}/(k_{f_{C}} \cdot \delta_{C}/2)}$$
 (4.12)

Taking, thermal conductivity,

 $k_{\mbox{f}_{\mbox{C}}} = 737.355 \ \mbox{kJ/m-h-°C}$  for Aluminium fin and substituting respective values in Eqs. (4.11 and 4.12), we have

 $\rm r_{\rm C}^{*}=5.025$  and  $\rm \Psi_{\rm C}=0.316$  and correspondingly  $\rm \emptyset_{\rm C}=0.93$ 

Substituting the respective values in Eqs. (4.8 and 4.10)

$$h_{red_c} = 17.63 \text{ kJ/m}^2 - h^\circ \text{C} \text{ and } U_{f_o} = 17.5845 \text{ kJ/m}^2 - h^\circ \text{C}$$

Heat transfer rate per unit length of the condenser is given by:

$$q_{c} = U_{f_{c}} \times (A_{f_{c}} + A_{o_{c}}) \times (T_{s_{c}} - T_{am})$$
 (4.13)

Substituting the respective values in Eq. (4.13) we have,

$$q_c = 17.5845 \times (0.76495 + 0.050878) \times (47 - 41.3)$$
  
= 81.7718 kJ/h

Since the heat rejected from the condenser is  $\dot{\Omega}_{_{\mbox{\scriptsize C}}}$  = 656 kJ/h

the length of the finned copper tube required for the complete condensation of the water vapour is:

$$L_c = Q_c/q_c = 8.0 \text{ m}$$

Now, this length of tube has to be arranged such as to have a compact volume of the condenser. This can be arranged in two columns and six rows, so that the condenser length is equal to 0.52 m and the rest tube is used in eleven bends. The assumed 72 fins per m length are to be mounted on 0.78 m length. Hence, the spacing to be kept is  $10.83 \times 10^{-3}$  m.

#### 4.2 Absorber Design

The main function of the absorber is to cool and totally condense the refrigerant vapour coming from the evaporator. The strong LiBr - water solution becomes a weak solution. This equipment is also called as absorber-condenser. In the simplest [31] absorber condenser design, the refrigerant vapours and the absorbent liquid are fed through an efficient mixer to the top of the heat exchanger and withdrawn from the bottom as a complete condensed liquid at, or slightly below, its saturation temperature. The most efficient heat transfer arrangement for this purpose is with the vapour liquid mixture flowing downward in a single pass inside vertical tubes surrounded by cooling water. This would be much costlier arrangement

due to high cost of maintenance for a small unit like the present one.

In the air cooled absorber-condenser the vapour liquid mixture flows inside a series of finned tubes, the air is blown across the tube bank and heat due to absorption of water vapour gets rejected. In its design, the conventional methods for calculating heat transfer and pressure drops for condensation inside tubes may be used. The specific correlations are decided according to vertical or horizontal orientation of the tube.

In the present work air cooled absorber has been selected due to low maintenance cost and to eliminate the use of electricity as in many rural areas electricity is not available. The strong LiBr - water solution and the refrigerant vapour enter the top of the absorber, rendering parallel - two phase flow inside the tube. The LiBr solution which flows inside finned tubes, gets cooled due to natural correction. As the vapour gets absorbed during the process it is similar to two phase condensation.

The design of the absorber follows the same procedure as in case of condenser design. Rewriting the equations for the absorber side.

Heat transfer coefficient inside horizontal tube.

$$h_{i_a} = 0.555 \left[ \frac{\rho_{1_a} \cdot (\rho_{1_a} - \rho_{v_a})_g k_{1_a}^3 \cdot h_{1v_a}}{\mu_{1_a} d. \frac{(T_{s_a} - T_{w_i})}{\sigma_{1_a}}} \right]^{0.25}$$
(4.14)

The properties of LiBr-water solution [32] and water vapour at the designed conditions are given in table 4.3. The tube and fin dimensions selected are given in table 4.4.

Table 4.3

Properties of LiBr-H2O solution and water vapour

LiBr-H <sub>2</sub> O solution at $61.4\%$ conc. and T = $47^{\circ}$ C					Sat. water vapour at T = 47°C		
$^{\rho}$ l <sub>a</sub> $(kg/m^3)$	kla (kJ/m-h-°C)	μ <sub>la</sub> (kg/m-h				μ <sub>V a</sub> (at 5°C) a )(kg/m-h)	
17 30	1,686	18.72	1.84	0.0705	2391.2	0.0324	

Table 4.4

Tube and fin dimensions for the absorber

Tube dimensions	×	Fin d	imensions
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	ī <sub>a</sub> (m) x10 <sup>-2</sup>	r <sub>a</sub> (m) x10 <sup>-2</sup>	6 <sub>a</sub> (m) x10-3
1.508 1.968 0.754 0.984 0.23	3.374	4.895	0.58

The Reynolds number is given by:

$$Re_{a} = \frac{d_{1a} G_{va}}{\mu_{va}} = \frac{d_{1a} x(m_{6} / (\frac{\pi}{4} d_{1a}^{2}))}{\mu_{va}} = 677.54 < 35,000$$

Similarly,

$$h'_{1v_a} = h_{1v_a} + 0.68 C_{p_L} (T_{s_a} - T_{w_{i_a}})$$
 (4.15)

Again the unknown temperatures  $T_{w_i}$ ,  $T_{w_o}$ ,  $T_f$  are evaluated using interative method, hence  $h_i$ ,  $h_o$  and  $h_f$  are obtained. Substituting values from tables (4.3 and 4.4) in Eqs.(4.14 and 4.15), we have,

$$h_{i_a} = 4976.682 \left[ \frac{h_{i_a}}{47 - T_{w_{i_a}}} \right]^{0.25}$$
 (4.16)

and 
$$h_{\text{Iv}_{a}} = 2391.2 + 0.68 \times 1.84 \times (T_{s_{a}} - T_{w_{i_{a}}})$$
 (4.17)

Iteratively,

$$T_{W_{\mathbf{i}_a}} = 46.97^{\circ}C$$

$$h_{lv_a}' = 2391.2^{\circ} kJ/kg$$

and 
$$h_{i_a} = 83621 \text{ kJ/m}^2 - h - °C$$

Rewriting the Eqs. (4.6 and 4.7) for outer heat transfer coefficients across tubes and vertical plate fins respectively, we have

(a) 
$$h_{o_a} = 4.752 \left(\frac{\Delta T'}{d_o}\right)_a^{0.25} \text{ kJ/m}^2 - h - °C$$
 (4.18)

(b) 
$$h_{fa} = 5.112 \left(\frac{\Delta T''}{1}\right)^{0.25}_{a} \text{ kJ/m}^{2}-h^{\circ}C$$
 (4.19)  $(\Delta T')_{a} = (T_{w_{oa}} - T_{am})$ ;  $T_{am} = 41.3^{\circ}C$   $(\Delta T'')_{a} = (T_{fa} - T_{am})$ 

From iteration

$$T_{W_{O_a}} = 46.95$$
°C and  $T_{f_a} = 45.6$ °C

Therefore,

$$h_{oa} = 19.56 \text{ kJ/m}^2 - h - ^{\circ}\text{C} \text{ and } h_{fa} = 17.18 \text{ kJ/m}^2 - h - ^{\circ}\text{C}$$

Rewriting Eq. (4.8) for absorber side,

$$\frac{1}{U_{f_o}} = FF_a \left[ \frac{1}{h_{i_a}} + \frac{\delta'_a}{k_{w_a}} \right] + \frac{1}{h_{red_a}}$$
 (4.20)

$$FF_{a} = (A_{f_{a}} + A_{o_{a}}) / A_{i_{a}}$$

$$A_{f_{a}} = 2 \pi n_{a} [(r_{a} + \delta_{a}/2)^{2} - r_{o_{a}}^{2}]$$
(4.21)

$$A_{o_a} = 2 \pi L_a (1 - n_a \delta_a) r_{o_a}$$

$$A_{i_a} = 2 \pi r_{i_a} L_a$$

$$h_{red_a} = (h_{f_a} p_a A_{f_a} + h_{o_a} A_{o_a})/(A_{f_a} + A_{o_a})$$
.... (4.22)

Assuming,  $n_a=84/\text{unit}$  length and substituting the respective values in Eqs. (4.21 and 4.22), we have  $A_{fa}=1.2285~\text{m}^2$ ,  $A_{oa}=0.0588~\text{m}^2$ ,  $A_{ia}=0.0474~\text{m}^2$  and  $FF_{a}=27.$ 

The conductivity of mild steel (for tube),  $k_{\rm wa} = 193.0$  (at 47°C) and the conductivity of G.I. Sheet (for fin),  $k_{\rm fa} = 209.4$  (at 41.3°C) in kJ/m-h-°C.

Similarly, the parameters:

$$r_a^* = (r_a + \delta_a/2)/r_o_a$$
 (4.23)

$$\Psi_{a} = \bar{I}_{a} \sqrt{h_{f_{a}} / (k_{f_{a}} \cdot \delta_{a}/2)^{2}}$$
 (4.24)

Substituting respective values in Eqs. (4.21 and 4.22) we have,  $r^{\#}$  = 5.0 and  $\Psi_{a}$  = 0.512 and correspondingly  $\emptyset_{a}$  = 0.8.

Eq. (4.22) gives ,  $h_{red} = 14.0 \text{ kJ/m}^2-h^{-\circ}C$ 

Substituting values in Eq. (4.20) we have,

$$U_{f_0} = 13.9 \text{ kJ/m}^2 - h - ^{\circ}\text{C}$$

Since,

$$q_a = U_{f_{o_a}} A_{f_{o_a}} (T_{s_a} - T_{am})$$
 (4.25)

= 
$$13.9 \times 1.2285 (47 - 41.3) = 97.334 kJ/h$$

Now heat of absorption is found as  $\dot{\Omega}_a = 715$  kJ/h Hence, length of finned mild steel tube required is:

$$L_a = \frac{\dot{Q}_a}{q_a} = 715/97.334 = 7.35 \text{ m}$$

Total number of fins =  $7.35 \times 84 = 618$ It is assumed that the number of fins,  $n_a = 84/m$  length, in order to minimise the space occupied by the absorber, it may be arranged in different rows.

Let 6 m length tube be finned and rest be used in bends etc.

Thus, 618 fins are to be mounted on 6 m tube,

i.e., On 1 m tube, fins mounted = 103 with spacing

approximately 9.7 mm

### 4.3 Evaporator Design

As the refrigerant enters the evaporator through the throttle valve, it starts boiling as shown in Fig. 5.1 [33]. It is assumed in this analysis that the pressure drop along the tube is small compared to the static pressure rendering constant saturation temperature in the evaporator. Heat transfer coefficient to single phase liquid under forced convection is given as [33]:

$$h_{i_{e}} = 0.17 \left[ k_{1}/d_{i} \right]_{e} \left[ G_{1}.d_{i}/\mu_{1} \right]_{e}^{0.33} \left[ C_{p_{1}}.\mu_{1}/k_{1} \right]_{e}^{0.43}$$

$$\left[ p_{r_{1}} / p_{r_{w}} \right]_{e}^{0.25} \left[ d_{i}^{3} \rho_{1}^{2} g \beta \Delta T /\mu_{1}^{2} \right]_{e}^{0.1}$$

$$(4.26)$$

This relationship is valid for heating in vertical upflow or cooling in vertical downflow for L/d > 50 and  $\left[\frac{\text{Gd}}{\mu_1}\right] < 2000$ . It also accounts for the influence of free convection. The properties of saturated liquid at  $T_e = 5^{\circ}\text{C}$  are given in table 4.5.

Table 4.5

<sup>ρ</sup> 1 <sub>e</sub> (kg/m <sup>3</sup> )	k <sub>l</sub> e (kJ/m-h-°C)	μ <sub>1</sub> e (kg/m-h)	Cple (kJ/kg°C)	d <sub>ie</sub> (m) x 10 <sup>-2</sup>	d <sub>o</sub> e (m) <sub>2</sub> x 10 <sup>-2</sup>
999.9	2.08	6.48	4.19	1.268	1.608

where,

$$G_{l_e} = m_6/(\frac{\pi}{4} d_{l_e}^2)$$

$$\beta = 1/(273.15 + 5) \text{ and } \Delta T = (T_{w_{l_e}} - T_{s_e})$$

Substituting the respective values  $G_{1e} = 2058.95 \text{ kg/m}^2\text{-h}$  and  $\beta = 3.6 \times 10^{-3}$ .

For small diameter tubes, fluid properties at the wall surface are approximately same as at the bulk stream, so that  $p_{r_s}/p_{r_{s,r}}=1.0$ .

T and T are the unknown temperatures which are calculated iteratively, hence,  $h_i$  and  $h_o$ . Substituting the respective values in Eq. (4.26)we have,

$$h_{i_0} = 362.84 (\Delta T)^{0.1}$$
 (4.27)

By iteration,  $T_{w_i} = 7.519^{\circ}\text{C}$  and  $h_i = 397.96 \text{ kJ/m}^2-h-^{\circ}\text{C}$ 

Heat transfer coefficient for helical coil [34],

$$h_{i_{e_{x}}} = h_{i_{e}} \times 1.127 = 448.5 \text{ kJ/m}^{2}-h^{\circ}C$$

Outside tube heat transfer coefficient by natural convection is given by [14] as:

Nu = 0.53 [Gr · Pr]<sup>0.25</sup> (4.28)  
for Gr.Pr = 
$$10^4$$
 to  $10^9$ 

or
$$h_{e} = \begin{bmatrix} \frac{k_{o}}{d_{o}} \end{bmatrix} \times 0.53 \times \begin{bmatrix} \frac{g \beta \Delta T'_{e} d_{o}^{3}}{v_{e}^{2}} \times Pr \end{bmatrix}^{0.25}$$

$$(4.29)$$

where, 
$$\Delta T'_e = (T_o - T_w)$$

Properties of water at 8°C taken from [24] are tabulated in table 4.6.

Table 4.6 Properties of water at  $T_{\infty} = 8^{\circ}C$ 

k °e (kJ/m-h-°C)	(m <sup>2</sup> /h) x 10 <sup>-3</sup>	Pr	β
2.05	5.3	10.97	( <del>273.15+8)</del>

Substituting values in Eq. (4.29) we have,

$$h_{e} = 1979.69 (\Delta T_{e})^{0.25}$$
 (4.30)

For helical coil

$$h_{o} = 2231.1106 (\Delta T_{e}')^{0.25}$$
 (4.31)

By iteration, 
$$T_{W_0} = 7.52^{\circ}C$$
, and  $h_0 = 1857.08 \text{ kJ/m}^2-h-^{\circ}C$ 

Overall heat transfer coefficient is given by

$$\frac{1}{U_{o_{e}}} = \frac{A_{o_{e}}}{A_{i_{e}}} \times \left[ \frac{1}{h_{i_{e}}} + \frac{\delta'_{e}}{k_{w_{e}}} \right] + \frac{1}{h_{o_{e}}}$$
 (4.32)

The inner and outer areas per unit length using table 4.5, are:

$$A_{i_e} = \pi d_{i_e} L_e = 0.0398 m^2$$
 $A_{o_e} = \pi d_{o_e} L_e = 0.051 m^2$ 

Since  $\delta'_e = 1.7 \times 10^{-3}$  m and the thermal conductivity of copper,  $k_{W_e} = 1392.6 \text{ kJ/m-h-°C}$ 

Substituting the respective values in Eq. (4.32) we have,

$$U_{o} = 296.956 \text{ kJ/m}^2 - h - °C$$

Total refrigeration load on the evaporator,

$$\dot{Q}_{a} = 600 \text{ kJ/h}$$

or

$$\dot{Q}_{e} = U_{o_{e}} \times \pi \times d_{o_{e}} \times L_{e} \times (\Delta T_{e})$$

$$\Delta T_{e} = (T_{o_{e}} - T_{s_{e}})$$
(4.33)

Substituting respective values, length of the evaporator tube,  $L_{\rm e}$  = 13 m.

For the design of a refrigerator for a certain value of  $\hat{Q}_e$ , finned coil can be used. In the design of it  $h_i$  be calculated from Eq. (4.26) and  $h_o$  and  $h_f$  from Eqs. (4.6 and 4.7), and the same procedure may be adopted

as in the design of the condenser and the absorber. If the water is to be cooled at 10°C, then the properties of water at 10°C would yield,

$$h_{e_{x}} = 8041.044 \text{ kJ/m}^2 - h - c$$

The value of  $h_{i}$  and other dimensions remaining same,  $e_{x}$ 

$$U_{oe} = 338.483 \text{ kJ/m}^2 - \text{h}^{\circ}\text{C}$$

and the length of the evaporator tube,

$$L_{e} = 11.0 \text{ m}$$

### 4.4 Generator Design

The heat transfer to generator renders the release of superheated water vapour. The generator is designed for operating pressure of 80 mm of  $H_g$  and  $T_{g_0}$  of 116°C. For this  $x_1$  = 61.4%,  $x_3$  = 0,  $x_7$  = 69%. For  $m_1$  = 1 kg/h

$$m_3 = 0.11 \text{ kg/h}$$

The change in quality of vapour in the generator is found to be

$$\bar{x} = \frac{0 + m}{2} = 0.055$$

From the flow pattern as shown in Figures (1.2 and 1.4) in [33] it is seen that the flow is bubbly flow for this quality of vapour.

The relation for boiling with bubbles [14] can be written as

$$Nu = \frac{1}{C_{s_f}} (Re)^{0.667} (Pr)^{-0.7}$$
 (4.34)

where,

Nu = 
$$(h_i D_b/k_1)_g$$
, Pr =  $(\mu_1 C_{p_1}/k_1)_g$ ,

Re =  $(\frac{Q_g}{A h_{lv}}, \frac{D_b}{\mu_1})_g$  and  $D_b = \sqrt{\frac{\sigma_t}{(\nu_1 - \nu_v)_g}}$ 

(4.35)

Whenever possible it is recommended [33,35] that a pool boiling experiment be carried out to determine the value of  $C_{\rm sf}$  applicable to the particular conditions of interest. In the absence of such information a value of  $C_{\rm sf}$  = 0.013 may be used as a first approximation. The value of  $C_{\rm sf}$  for the water-polished stainless steel [27] is 0.0132.

The values of the various parameters taken from [14,24,32] at 61.4% concentration and generator temperature,  $T_g = 116^{\circ}\text{C}$  are given in table 4.7

Table 4.7

Property values of LiBr-H<sub>2</sub>O and water vapour (at 116°C and 64.1%)

σ <sub>t</sub>	k <sub>1</sub> g	$\mu_{1_g}$	c <sub>p</sub> lg	<sup>6</sup> lg	$^{\rho}v_{g}$
$\frac{(N/m)}{x}$ 10 <sup>-3</sup>	(kJ/m-h-°C)	(kg/m-h)	(kJ/kg°C)	(kg/m <sup>3</sup> )	$(kg/m^3)$
68.4	1.78	7.2	1.84	1.68x10 <sup>3</sup>	0.056

where,

$$\mathcal{T}_{t}$$
 = surface tension of water  $\mathcal{F}_{v_g}$  = density of water vapour

and

$$c_{p_{l_q}}$$
,  $\mu_{l_q}$ ,  $k_{l_q}$ ,  $p_{l_q}$  are respectively

specific heat, dynamic viscosity, thermal conductivity and density of LiBr-H,0 solution.

Substituting the respective values, in Eq. (4.34) we have,

$$D_{b} = \sqrt{\frac{68.134 \times 10^{-3}}{(1.68 \times 10^{3} - 0.056^{\circ}) \times 9.81}} = 2.038 \times 10^{-3} \text{ m}$$

For  $\dot{Q}_{\rm g}$  = 830 kJ/h and tube dimensions as in table 4.8 we have,

$$A_{o_{G}} = \frac{\pi}{4} (2.068 \times 10^{-2})^2 = 3.3588 \times 10^{-1} \text{ m}^2$$

and

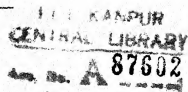
$$h_{1v}' = 2391.2 + 0.68 \times C_{p_{1_g}} \times (T_{w_{0_g}}^{-T} s_g)$$

$$= 2391.2 + 0.68 \times 1.839 \times (120-116)$$

$$= 2396.2 \text{ kJ/kg.}$$

Table 4.8

Dimensions of	of generator t	ube
d <sub>i</sub> g (m) x 10 <sup>-2</sup>	d <sub>o</sub> g (m) x 10 <sup>-2</sup>	(m) x 10 <sup>-3</sup>
1.68	2.07	1.96



Substituting values in Eq. (4.35) we have,

Re = 0.292 and

Pr = 7.43

Therefore, from Eq. (4.34) taking  $C_{s_f} = 0.0132$  and putting respective values:

$$h_{ig} = 7162.15 \text{ kJ/m}^2 - h - ^{\circ}C$$

To heat the weak solution flowing through the generator heat is supplied by burning biogas inside the generator outer casing. The appropriate size of the generator has been calculated by iteration.

The first approximation assumes the length of the enclosure carrying a single tube (as an annulus) having  $L_g = l_g = 0.305$  m. Let the inner diameter of the enclosure,  $D_i = 0.13$  m and outer diameter of the tube,  $d_0 = 0.02068$  m. The shape factor was calculated as per [36]. It is found that  $F_{1-2} = 1.4$  for the above dimensions. Here the enclosure is considered as body 2 and the tube as body 1. Heat transfer coefficient due to radiation is given by:

$$h_{r_g} = \frac{F_{1-2} \times \varepsilon_r \times \sigma \times \left[ \left( T_{g_{aV}} + 273.15 \right)^4 - \left( T_{g_o} + 273.15 \right)^4 \right]}{\left( T_{g_{aV}} - T_{g_o} \right)}$$

where,  $\epsilon_r$  = emmissivity (approx. 1.0 for a black surface)  $\sigma = \text{Stephan Boltzman constant} \ (= 20.42 \times 10^{-8} \ \text{kJ/m}^2 - \text{h-K}^4)$ 

lg = length of the enclosure, m

 $T_{gav}$  = average temperature of the flue gas in the enclosure

T<sub>g</sub> = outer surface temperature of the tube.

If the flue gas is entering the enclosure at 166°C and leaving it at 145°C then,  $T_{g_{av}} = 156$ °C.

Substituting the respective values in Eq. (4.36) we have,

$$h_{r_g} = 79.4 \text{ kJ/m}^2 - h - ^{\circ}C$$

Heat transfer coefficient due to free convection of air given as

$$h_{c_g} = 4.752 \left( \Delta T_{m_g} / d_{o_g} \right)^{0.25}$$
 (4.37)

Now LMTD,

$$\Delta T_{\text{mg}} = \frac{166 - 145}{\ln(\frac{166 - 117}{145 - 117})} = 37.53^{\circ}\text{C}$$

Substituting values in Eq. 4.37 we have,

$$h_{c_g} = 31.02 \text{ kJ/m}^2 - h - ^{\circ}C$$

Total outer heat transfer coefficient,

$$h_{o_{g}} = h_{r_{g}} + h_{c_{g}} = 110.407 \text{ kJ/m}^{2} - h^{\circ} C$$

$$\frac{1}{U_{o_{g}}} = \frac{d_{o_{g}}}{d_{i_{g}}} \left[ \frac{1}{h_{i_{g}}} + \frac{\delta'_{g}}{k_{g}} \right] + \frac{1}{h_{o_{g}}} \qquad \cdots \qquad (4.38)$$

Substituting the values in Eq. (4.38) we have,

$$U_{o_{G}} = 108.21 \text{ kJ/m}^2 - h - ^{\circ}C$$

Using, 
$$\dot{Q}_g = A_{o_g}$$
  $U_{o_g}$   $\Delta T_{m_g}$ , we get 
$$A_{o_g} = 0.204 \text{ m}^2$$
 or 
$$L_g = 3.14 \text{ m}$$

The length of the generator tube,  $L_g$  comes out to be 3.14 m which is more than the assumed value. For the next approximation four tubes were taken having the same dimensions kept within the same enclosure.

The shape factor,  $\mathbf{F}_{1-2}$  , for this geometry is found graphically as shown in Fig.(4.1)and can be written as follows:

$$F_{1-2} = \frac{1}{A_1} J_{A_1} J_{A_2} \frac{\cos \theta_1 \cos \theta_2}{\pi s^2} d A_1 d A_2$$
 (4.39)

Since,  $\theta_1 = 0$  and the elemental area  $dA_1$  of  $A_1$  are equal segments, therefore,

$$F_{1-2} = \frac{1}{A_1} \left[ dA_1 \sum_{j=1}^{8} \frac{\cos \theta_2}{\pi s_j^2} dA_2 \right] + \frac{1}{A_1'} \left[ dA_1' \sum_{j=9}^{10} \frac{\cos \theta_2}{\pi s_j^2} dA_2 \right]$$

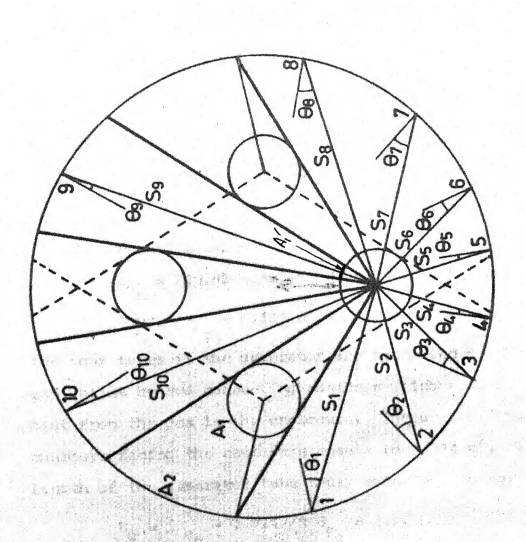
Substituting the values of 8 and S we have from Fig(4.1)

$$F_{1-2} = \frac{1}{A_1} \left[ 2 \times \frac{dA_1}{\pi} \left\{ \frac{\cos(29)}{(0.056)^2} \times 0.044 + \frac{\cos(28)}{(0.039)^2} \times 0.03 + \frac{\cos(20)}{(0.027)^2} \times 0.022 + \frac{\cos(8)}{(0.023)^2} \times 0.02 \right\} \times 0.305 \right]$$

$$+\frac{1}{A_1}\left[2\times\frac{dA_1}{\pi}\left\{\frac{\cos(11)}{(0.084)^2}\right\}\times 0.305\right]$$

From the Fig. (4.1),  $dA_1/A_1 = 1/8$  and  $dA_1/A_1' = 1$ 

$$\Theta_1 = \Theta_6 = 29^\circ$$
 $\Theta_2 = \Theta_7 = 28^\circ$ 
 $\Theta_3 = \Theta_6 = 20^\circ$ 
 $\Theta_4 = \Theta_5 = 8^\circ$ 
 $\Theta_9 = \Theta_{10} = 11^\circ$ 
 $S_1 = S_8 = 0.056 m$ 
 $S_2 = S_7 = 0.039$ 
 $S_3 = S_6 = 0.027$ 
 $S_4 = S_5 = 0.023$ 
 $S_5 = S_5 = 0.023$ 



generator Fig. 4.1 Radiative heat exchange in the

Ratio 1:1

Hence,  $F_{1-2} = 3.4$ 

It is seen from the Fig. (4.1) that the areas  $A_2$  and  $A_2'$  of the enclosure seen by a single tube is also seen by the adjuscent two tubes, due to which net effect on a single tube is about 25% of the calculated value of the shape factor.

i.e., 
$$F_{1-2} = 0.25 \times 3.4 = 0.85$$

Hence, the heat transfer coefficient within the enclosure carrying four tubes is given by:

$$h_{r_g} = 4F_{1-2} \times \varepsilon_r \times \sigma_x = \frac{\left[ (T_{g_{av}} + 273.15)^4 - (T_{g_o} + 273.15)^4 - (T_{g_o} + 273.15)^4 - (T_{g_o} + 273.15)^4 \right]}{(T_{g_{av}} - T_{g_o})}$$

Substituting the respective values we have,

$$h_{r_{cr}} = 192.8 \text{ kJ/m}^2 - h - ^{\circ}C$$

$$h_{o_g} = 223.82 \text{ kJ/m}^2-h-^{\circ}\text{C}$$
 and  $U_{o_g} = 214.982 \text{ kJ/m}^2-h-^{\circ}\text{C}$ .  
Thus,  $A_{o_g} = 0.103 \text{ m}^2$ 

The four tubes in the generator are connected to the inlet and outlet by two connecting chambers which also receive heat from the gas in the enclosure. The area of the two chambers facing the enclosure equals to 0.024 m<sup>2</sup>. The length of the generator tube thus, comes out to be:

$$L_g = (A_{o_g} - 0.024)/\pi d_{o_g} = 1.22 \text{ m}$$

This value is same as the assumed value.

Maximum possible heat transfer,

$$\dot{Q}_{p_{\text{max}}} = (\dot{m} c)_{\text{min.}} (T_{g} - T_{a})$$
 (4.43)

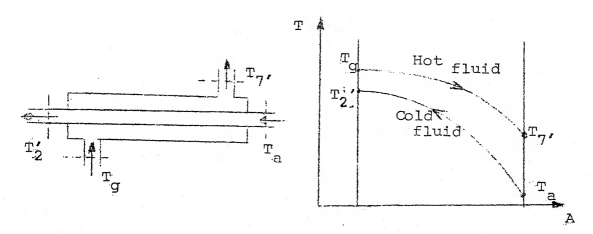


Fig. 4.2: Temperature distribution in the preheater.

The specific heats of LiBr-H<sub>2</sub>O solution at different concentrations are listed in table 4.9.

Table 4.9 Specific heats of LiBr-H $_2$ O solution along with mass flow rates

C <sub>p<sub>L</sub>a</sub> kJ/kg°C	<sup>™</sup> 1 kg/h	C <sub>pL</sub> · m1 kJ/h-°C	<sup>C</sup> p <sub>L</sub> g k∴/kg°C	<sup>fh</sup> 7 kg/h	C <sub>p</sub> . <sup>fl</sup> 7 g kJ/h-°C
1.839	2.301	4.34	1.73 (69g)	2.361	3.64

From table 4.9,  $\hat{m}_7$   $C_{p_L}$  is minimum.

The unknown temperatures  $T_2'$  and  $T_7'$  are determined using effectiveness value,  $\epsilon_2 = 0.75$ 

i.e. 
$$\varepsilon = \{ \tilde{m}_1 \, C_{p_L} \, (T_2' - T_a) \} / \{ \tilde{m}_7 \cdot C_{p_L} \, (T_g - T_a) \}$$
... (4.44)

Also,

Since  $T_{\alpha} = 116^{\circ}C$  and  $T_{a} = 47^{\circ}C$ ,

substituting the respective values in Eqs. (4.44 and 4.45) we have,

$$T_2' = 90.34^{\circ}C$$

$$T_7' = 64.25^{\circ}C$$

The log mean temperature difference on substitution of respective values is given as:

$$\Delta T_{m} = \frac{(T_{g} - T_{2}') - (T_{7}' - T_{a})}{1n \frac{T_{g} - T_{2}'}{T_{7}' - T_{a}}} = 21.2^{\circ}C$$

In fluction properties are taken at average temperatures for evaluation of the heat transfer coefficient. Annulus (he , strong solution),  $T_h = (T_g + T_7')/2 = (116 + 64.25)/2 = 90.12°C$ . Inner tube (cold, weak solution),

$$T_C = \frac{T_a + T_2'}{2} = \frac{47 + 90.34}{2} = 68.67$$
°C

Properties of LiBr-H<sub>2</sub>O solution at different conditions are given in table 4.10.

Table 4.10

Properties of LiBr-H20 solution at different conditions

Annull	ar side 0.12°C)	and blue			
kJ/m-h-°C	μ <sub>o</sub> p	<sup>P</sup> .o <sub>p</sub> (kg/m <sup>3</sup> )	k <sub>i</sub> pkJ/m-h-°C	μ <sub>i</sub> p	ρ p (kg/m <sup>3</sup> )
1.75	15.12	1858	1.748	12.24	1715

Assuming the following dimensions:

Annulus: 
$$D_{ip} = 2.841 \times 10^{-2} \text{ m}$$
;  $d_{op} = 2.068 \times 10^{-2} \text{ m}$   
Area,  $A_{ap} = \frac{\pi}{4} (D_{ip}^2 - d_{op}^2) = 2.98 \times 10^{-4} \text{ m}^2$ 

Equivalent diameter, 
$$D_{\frac{1}{2}p}^2 - d_{0p}^2 = 1.336 \times 10^{-2} \text{ m}$$

$$D_{e_p} = \frac{D_{\frac{1}{2}p}}{D_{\frac{1}{2}p}} = 1.336 \times 10^{-2} \text{ m}$$

$$Tube, d_{\frac{1}{2}p} = 1.676 \times 10^{-2} \text{ m}, d_{0p} = 2.068 \times 10^{-2} \text{ m}$$

$$A_{\frac{1}{2}p} = \frac{\pi}{4} d_{\frac{1}{2}p}^2 = 2.206 \times 10^{-4} \text{ m}^2$$

Carrying out the calculations for the annulus and the tube separately we have,

Annulus:  
Re = 
$$\frac{G_{\text{op}}}{\mu_{\text{op}}} = \frac{m_7}{A_{\text{ap}}} \times \frac{D_{\text{ep}}}{\mu_{\text{op}}} = \frac{2.101 \times 1.336 \times 10^{-2}}{2.98 \times 10^{-4} \times 15.12} = 6.23$$

$$Pr_{o_{p}} = \frac{\mu_{o_{p}} \cdot C_{p}}{k_{o_{p}}} = \frac{15.12 \times 1.73}{1.75} = 14.944$$

$$G_{r_{o_{p}}} = \frac{g^{\beta_{o_{p}} \wedge T_{m}} \cdot D_{e_{p}}^{3} \rho^{2}_{o_{p}}}{\mu_{o_{p}}^{2}} = 9.31 \times (3600)^{2} \times \frac{1}{\cancel{90.02+273.15}}$$

$$\times \frac{21.2 \times (1.336 \times 10^{-2})^{3} \times (1858)^{2}}{(15.12)^{2}} = 2.672 \times 10^{5}$$

Assuming length of preheater as 1 m

$$Gz_{op} = Re_{op} \cdot Pr_{op} \cdot \frac{D_{ep}}{L_{p}} = \frac{6.23 \times 14.944 \times 1.336 \times 10^{-2}}{1.0} = 1.244$$

 $\mu_h = \mu_m$ Let

Substituting these values in Eq. (4.40) we have heat transfer coefficient on the annulus side:

$$h_{op} = 402.144 \text{ kJ/m}^2 - h - C$$

Tube:

$$Re_{i_{p}} = \frac{G_{i_{p}}^{d_{i_{p}}} G_{i_{p}}}{\mu_{i_{p}}} = \frac{M_{1}}{A_{i_{p}}} \times \frac{G_{i_{p}}^{d_{i_{p}}}}{\mu_{i_{p}}} = \frac{2.361}{2.206 \times 10^{-4}} \times \frac{1.676 \times 10^{-2}}{12.24}$$

$$= 14.655$$

$$Pr_{i_{p}} = \frac{\mu_{i_{p}}^{C_{p_{i_{p}}}} G_{p_{i_{p}}}}{\mu_{i_{p}}} = \frac{12.24 \times 1.839}{1.748} = 12.876$$

$$Gr_{ip} = \frac{g \beta_{ip}^{\Delta T_{m}} \times d_{ip}^{3 \rho 2}}{\mu_{ip}^{2}} = 9.81 \times (3600)^{2} \times \frac{1}{(68.61+273.15)}$$

$$\times \frac{21.2 \times (1.676 \times 10^{-2})^3 \times (1715)^2}{(12.24)^2} = 7.29 \times 10^5$$

$$Gz_{ip} = Re_{ip} \cdot Pr_{ip} \cdot \frac{d_{ip}}{L_p} = 14.655 \times 12.876 \times \frac{1.676 \times 10^{-2}}{1.0}$$
  
= 3.1626 (L = 1.0 m)

Substituting the respective values in Eq. (4.40), heat transfer coefficient on the inner side:

$$h_{ip} = 307.54 \text{ kJ/m}^2 - h_o^{\circ}$$

The overall heat transfer coefficient in the preheater is:

$$\frac{1}{U_{o_p}} = \frac{D_{i_p}}{d_{i_p}} \cdot \frac{1}{h_{i_p}} + \frac{D_{i_p}}{2 k_{w_p}} \times \ln \left(\frac{d_{o_p}}{d_{i_p}}\right) + \frac{D_{i_p}}{2 k_{w_p}} \cdot \ln \left(\frac{D_{i_p}}{d_{o_p}}\right) + \frac{1}{h_{o_p}}$$
.... (4.46)

The conductivity of mild steel,

$$k_{\text{Wp}} = 193 \text{ kJ/m-h-}^{\circ}\text{C}$$

Substituting the respective values in Eq. (4.46) we have,

$$U_{op} = 124.47 \text{ kJ/m}^2 - h - ^{\circ}C$$

From Eq. (4.41)

$$\dot{Q}_{p} = 2.361 \times 1.839 \times 21.2 = 92.05$$

Using,

$$Q_{p} = U_{o_{p}} \cdot A_{o_{p}} \cdot \Delta_{m} \qquad (4.47)$$

and putting the respective values,

$$A_{o_p} = 0.035 \text{ m}^2$$
 and  $L_p = 0.84 \text{ m}$ 

Thus, the value of L taken equal to 1.0 m is correct.

# 4.6 Fabrication and Assembly

Table 4.11 shows the specifications of the different components of the system. Fig. (4.3) shows the schematic view of the set-up. The condenser, absorber, generator and preheater were fabricated by 'Kalyan Kooling Corporation', Kanpur: As the water vapour condenses in the condenser, it was fabricated from copper having Aluminium fins. The absorber was fabricated from mild steel tube having G.I. sheet fins as shown in Fig. (4.4). The generator was fabricated from mild steel tube of 16.8 mm inner diameter. Four tubes were welded in 130 mm inner diameter pipe. The exit of the generator has two outlets : one for vapour supply to the condenser and the other for return of strong solution to the absorber via preheater. The evaporator is a helical coil copper tube fabricated from 16.08 mm outer diameter and is immersed in a water tank. Generator, heat exchanger, and the water tank were thermally insulated.

The assembly of the whole system is shown in Fig. (4.4). To measure the operating pressures in the generator and evaporator sides, inclined tube mercury manometers have been used. The connection between generator and condenser is done using glass tubes for visual aid. A drain valve is provided at the lowest level in the preheater. The charging line is provided on the

absorber side. The absorber is placed on adjustable jacks and the connections to the frame are made through polythene tubes. This is done for adjustment of pressure difference between the absorber and the generator. Thermocouples are provided to measure temperature in the evaporator and the generator.

Table 4.11 Specifications of the system components

Components Specifications	Conden- ser	Absor- ber	Evapora- tor	Gene- rator	Preheater
Heat transfer rate, Q (kJ/h)	57 <b>65 6</b> 38	715	600	830	_
Tube dimensions:					Tube:
Length (m)	8.0	7 4 35	13.00	1.22	1.0
$d_0 \times 10^{-2} \text{ (m)}$	1.67	1.968	1.608	2.073	2.068
$d_1 \times 10^{-2} \text{ (m)}$	1.43	1.508	1.268	1.68	1.676
Material	copper	M.S.	Copper	M.S.	M.S
Square Fin dimensions:					Annulus:
$\overline{1}$ x10 <sup>-2</sup> (m)	2.8675	3.374			$D_{i} = 2.841$ x $10^{-2}$ m
\$x10 <sup>-3</sup> (m)	0.42	0,58			$D_0 = 3.541$ x $10^{-2}$ m
Spacing (m) x 10 <sup>-3</sup>	10.83	9 ,70			Material: M.S.
Material	Al.	G.I. sheet			

The biogas plant has been fabricated from a cylindrical drum having a floating top as shown in Fig. (4.5). The inlet for charging and exit for discharge of disgested gobar are provided. The gas from the plant is supplied to generator burner through a G.I. Sheet. Wet-test metre is provided to get the flow rate of biogas. The manometer is provided in the supply line of the biogas to get the supply pressure.

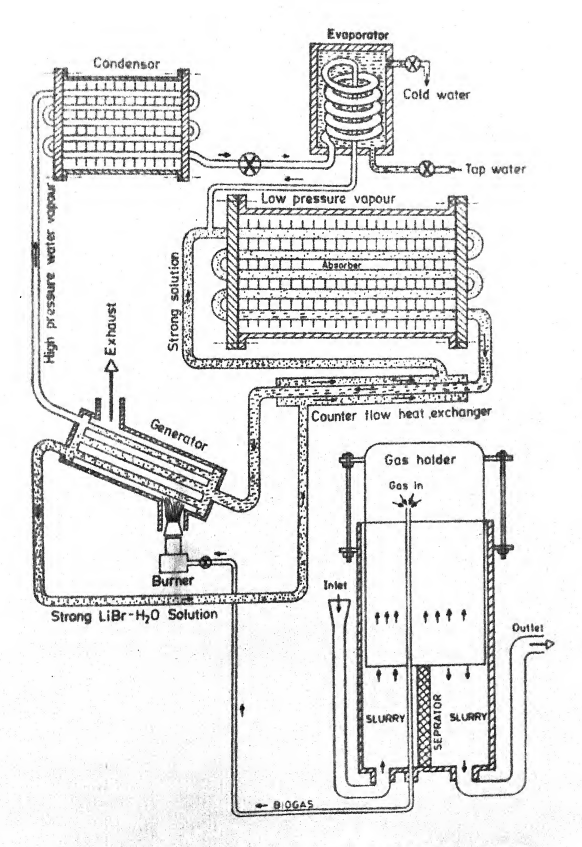


Fig. 4.3 Schematic view of the experimental set-up.



Fig. 4,4 Experimental set-up



Fig. 4.5 View of Gobar-Gas Plant

## CHAPTER - V

### RESULTS AND DISCUSSION

## 5.1 Analytical Results

COP (for  $P_g$  = 40, 50, 60, 70 and 80 mm of Hg and HE effectiveness,  $\epsilon$  = 0.75) and  $\Omega_{gas}$  with  $T_g$  are shown in Fig. (2.2) having concentration below 70%. Evidently the variation in COP with  $T_g$  is more dominant than the biogas energy,  $\Omega_{gas}$ .

The cost of biogas was computed for  $p_g=40$ , 50, 60, 70 and 80 mm of Hg having HE effectivenesses of 0.65, 0.75 and 0.85 for 0.7, 1.4 and 2.1 kW of cooling capacities. The life of the biogas plant was assumed as 15 years having 10% interest rate. Analysis of the system has been carried out with precooler (PC),  $\frac{\varepsilon}{1}$  and preheater (PH),  $\frac{\varepsilon}{2}$  both (i.e.,  $\frac{\varepsilon}{1} = \frac{\varepsilon}{2} = \frac{\varepsilon}{2}$ ) and without precooler (i.e.,  $\frac{\varepsilon}{1} = 0$ , using PH only). Figures (5.1 to 5.5) exhibit variation in cost with generator temperature for a given generator pressure and evaporator temperature for different tonnages.

# (i) Variation in cost with pressure

The cost variation with  $T_g$  for different pressures  $(p_g)$  show that the system operating at higher pressures are costlier than that at lower pressures. With the difference of 10 mm of Hg between two pressures, the cost variation is about 1.3%.

### (ii) Variation in cost with TR

Cost variation for a fixed generator pressure and effectiveness differs significantly for all values of TR Figs. (5.1 to 5.4) whereas, the  $T_{go}$  values for all TR are same as is evident from Eq. (2.33).

Figures (5.6 and 5.7) represent the optimum generator temperature and cost of biogas for various values of parameters.

# (iii) Variation in cost with HE effectiveness

For decrease in effectiveness for about 10% the cost increases for about 2% under optimum operating conditions. It shows a linear variation in cost with change in effectiveness within ranges of our study.

Analysis have been carried out for:

- (a) using precooler and preheater both (i.e.,  $\epsilon_1 = \epsilon_2 = \epsilon$ )
- (b) with preheater only (i.e.,  $\epsilon_1 = 0$ )
  The results for these two cases are shown in Fig. (5.8).

It is seen that the latter is costlier (about 1.5 to 2%) than the former. But the latter is generally allowed in small units in view of simplicity.

The multiplier shown in Fig. (5.9) can be used to determine  $T_{go}$  for evaporator temperatures other than 5°C. To do this one should get  $T_{go}$  from Figures (5.1 to 5.5) for a set of operating variables and evaporator temperature of 5°C. Then this  $T_{go}$  is multiplied by the multiplier to get the required value.

## 5.2 Experimental Set-up

The system components have been fabricated and assembled. The absorber is kept at about 80 cm above the generator. Inclined Hg-manometers are used to measure the generator and evaporator pressures. Water manometer and wet-test meter are used to get the pressure and the flow rate of the biogas, respectively. Most of the joints are brazed. Only the polythene tube connections to the absorber are clamped and sealed using epoxy. The system has been evacuated to 1 atmosphere. The experiment was not possible due to some minor leakages. After a few months of trial it has been realized that it would take longer time which became a contraint in the present time bound programme.

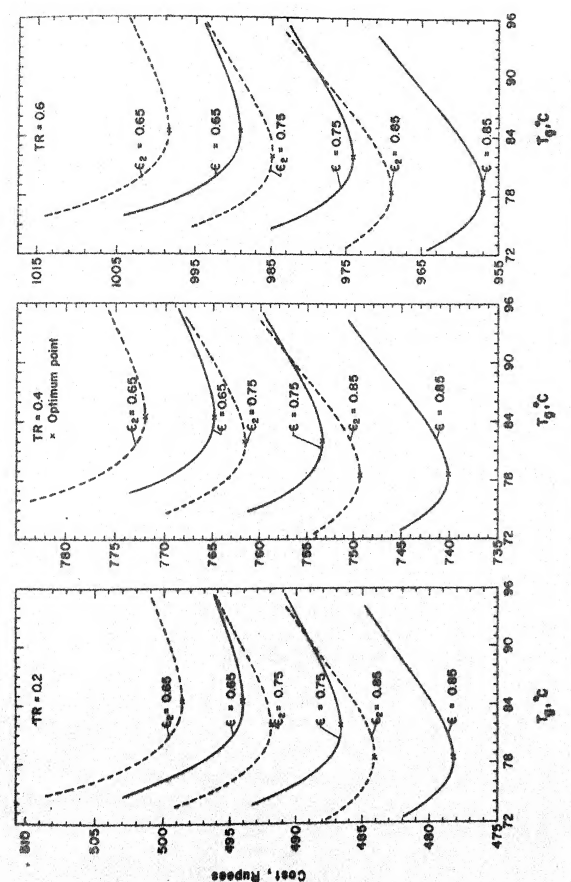
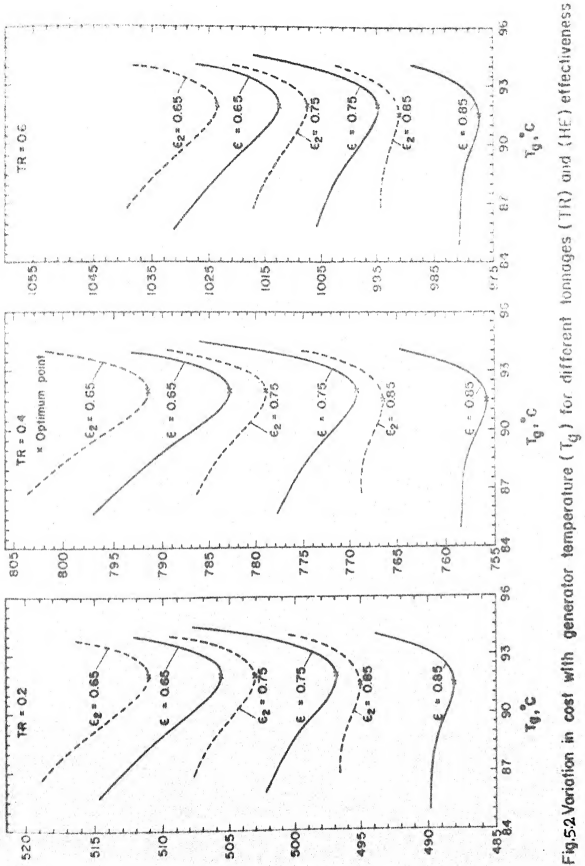


Fig.5.1 Variation in cost with generator temperature (Tg) for different tonnages (TR) and HE effectiveness (Pg = 40 mm of mercury and Te = 5°C)



Cost,

seedny

Pg = 50 mm of mercury and Te = 5 °C

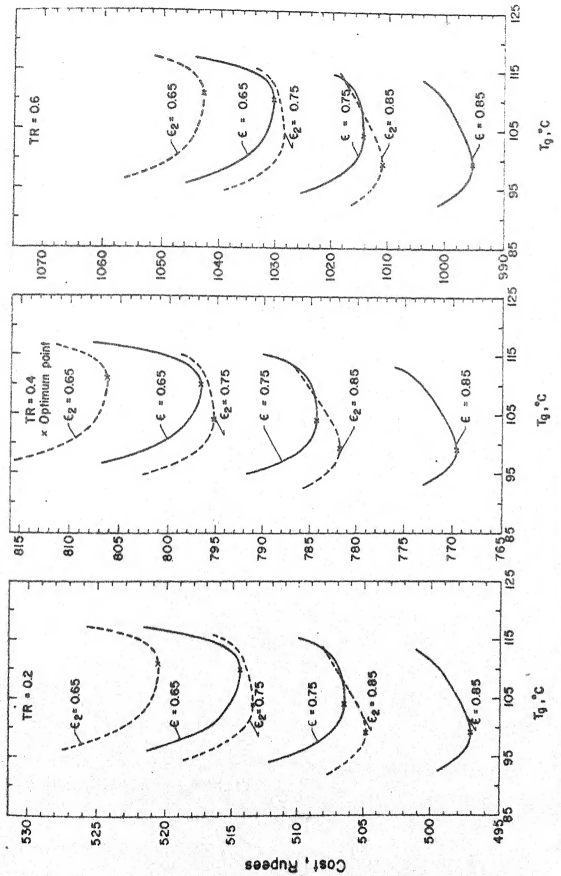


Fig.5.3 Variation in cost with generator temperature Tg , for different tonnages (TR) and HE diffectivnesses ( $p_g = 60 \, \text{mm}$  of mercury and  $T_e = 5^{\circ}C$ ).

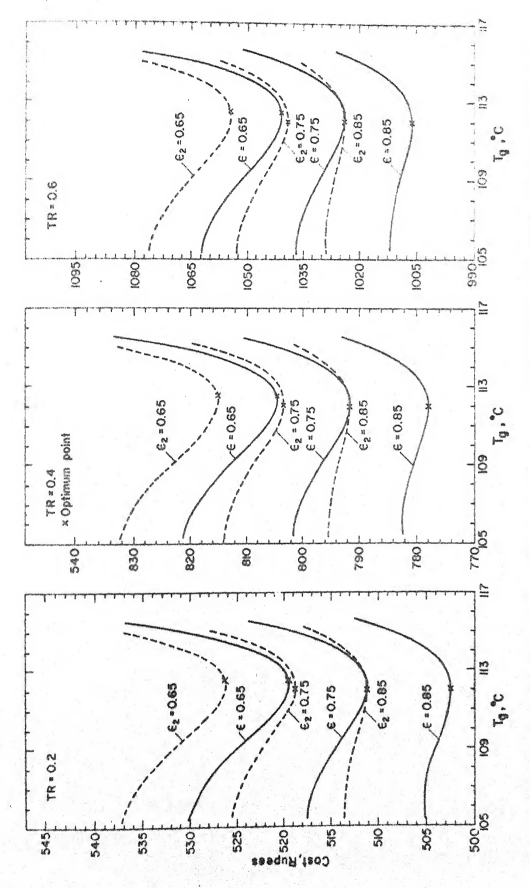


Fig.5.4 Variation in cost with generator temperature Tg., for different tonnages (TR) and HE effectiveness  $(p_g = 70 \text{ mm of mercury and } T_e = 5^{\circ}C)$ 

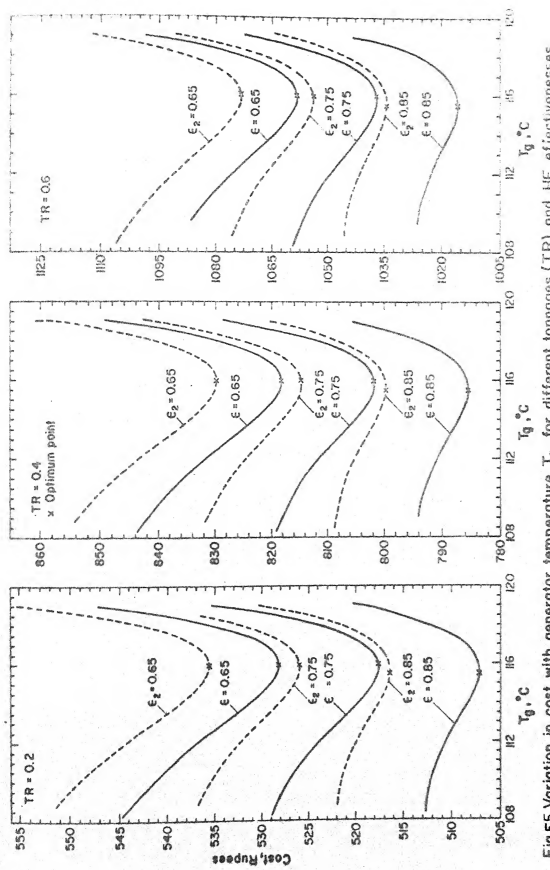


Fig.5:5 Variation in cost with generator temperature Tg, for different tonnages (TR) and HE effectivenesses (pg=80 mm of mercury and Te=5°C)

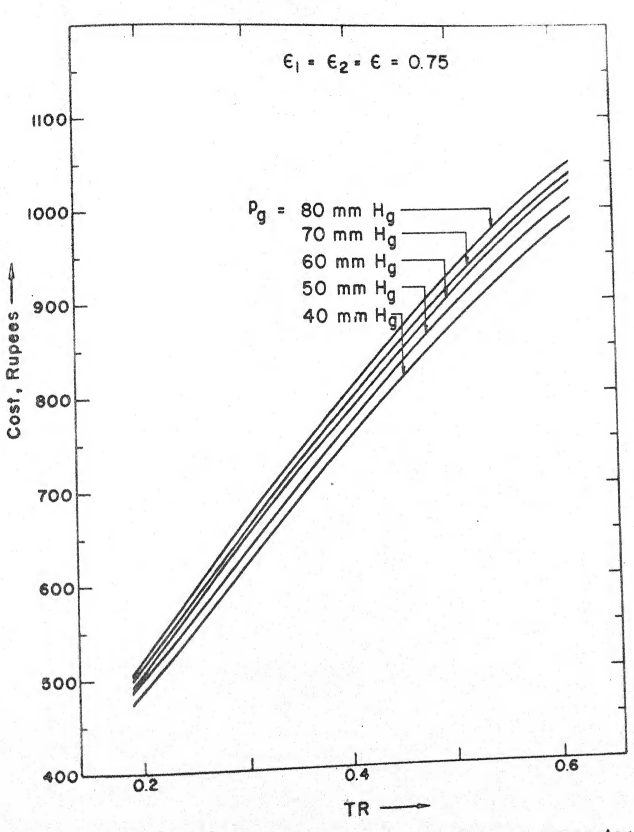
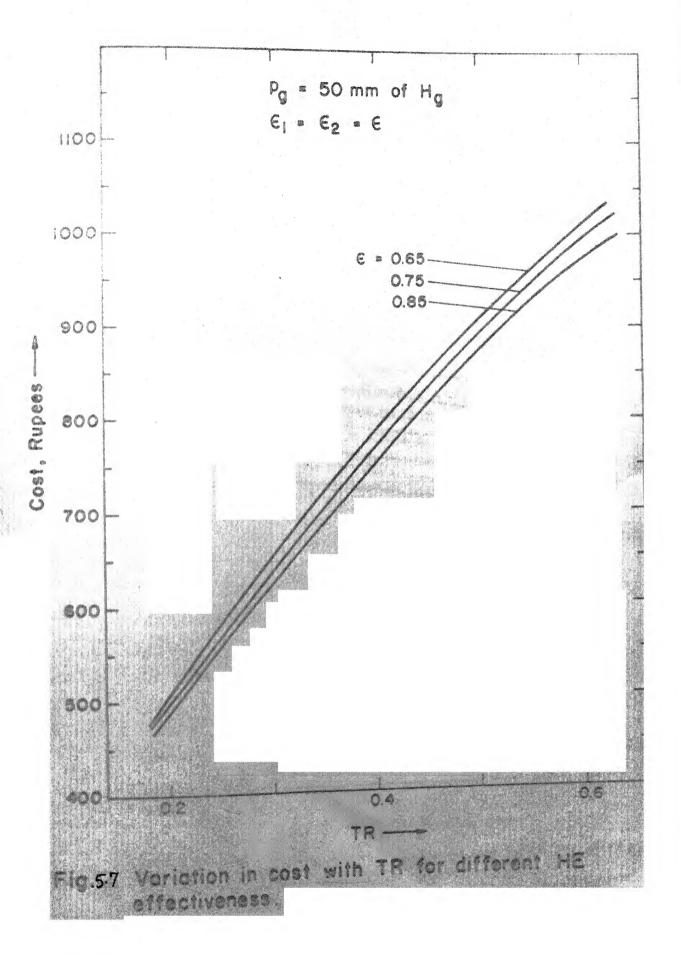


Fig.5.6 Variation in cost with TR for different generator pressures Pa.



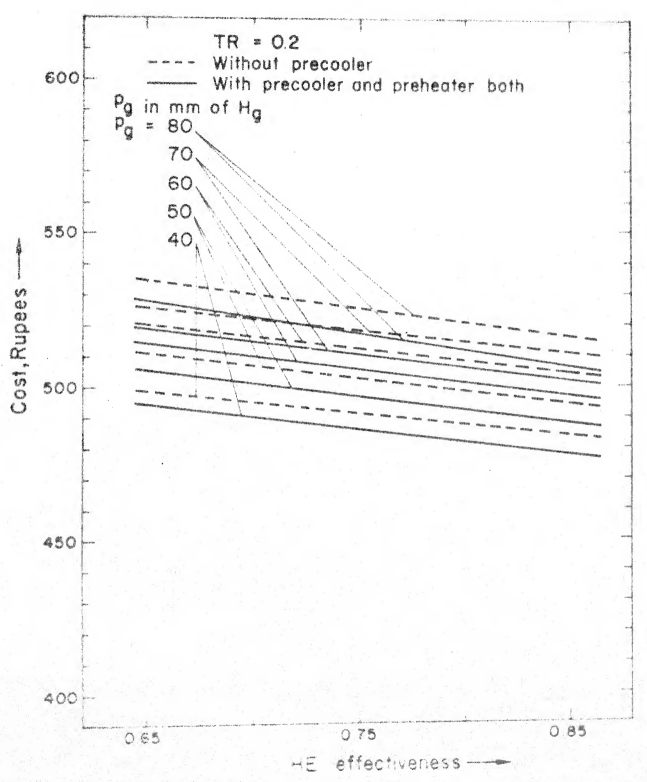


Fig. 5.8 Variation in cost with HE effectiveness for differen generator pressures p<sub>a</sub>.

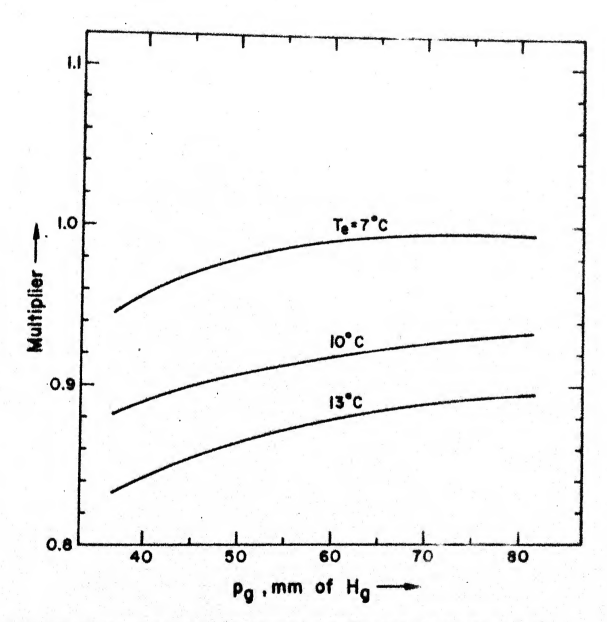


Fig.59 Multiplier for optimum generator temperature

The biogas plant was fed with 250 kg of slurry (equal weight of gobar + water) upto 0.23 m<sup>3</sup> volume of the drum initially. The recharging was started after 22 days of the initial charge, and was continued with 10 kg of slurry. The biogas generated from the plant was available at about 200 mm water head when heat was applied to the digester cylinder. Under normal winter temperature, the pressure of the biogas has been measured to be 100 mm of water.

# CHAPTER - VI

#### CONCLUSION

- 1. The optimum generator temperature has been computed for the LiBr-H2O absorption system using biogas energy for a wide range of operating conditions.
- 2. The biogas cost goes up by 1.3% for every 10 mm of rise in the generator pressure.
- 3. The optimum cost of the biogas for the system goes up by about 2% for every 10% decrease in effectivenesses of heat exchangers.
- 4. The cost of biog is goes up by 1.5 to 2% if the system operates without a precooler.
- 5. The multipliers have been presented for utilization of the present optimum results for evaporator temperatures other than 5°C.
- 6. To facilitate the user the optimum values of generator temperature are presented using capacity of refrigeration system as a variable for different operating variable.

- 7. The procedure is developed for design of various components of the vapour-absorption system.
- After fabrication of all designed components, the system has been assembled for experimental purpose. The provisions are made for adjustment of relative elevation between the generator and absorber. This enables us to conduct experiment at other operating conditions as well.
- 9. The biogas was generated and is ready for the use in the system.
- 10. Components for this refrigeration system when fabricated and assembled in the laboratory has the problem of leakage. This is more so because it has to operate at low pressures which is possible only if the system is completely evacuated. The leakage problem becomes more acute because, in order to instrument the system, number of gauges and temperature transducers had to be inserted at several points. Making it leakage free is one of the most time consuming job. The leakage will not be a problem if the sealing etc. is done more professionally.

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#### APPENDIX - A

# Thermodynamic Properties of the LiBr-H20 Solution and Water

From standard references [14,16,32] the various properties are listed below:

Properties (i) to (ix) are in F.P.S. units (h, Btu/lb, T. °F and P. psia).

- (i) Solution temperature: T = AT' + B
- (ii) Refrigerant temperature : T' = (T B)/A

(iii) 
$$A = -2.00755 + 0.16976 \times - (3.133362 E - 3) \times^{2} + (1.97668 E - 5) \times^{3}$$

(iv) 
$$B = 321.128 - 19.322 \times + 0.374382 \times^2 - (2.0637 E - 3) \times^3$$

(v) Vapour pressure : 
$$\log_{10} P = C + D/(T' + 459.72)$$
  
E/  $(T' + 459.72)^2$ 

where,

$$C = 6.21147$$
  $0 \ge T' \le 230 \text{ °F}$ 
 $D = -2886.373$   $40 \ge T \le 350 \text{ °C}$ 
 $E = -337269.46$   $45\% \ge x \le 70\%$ 

(vi) Enthalpy of the solution: 
$$h = A_1 + A_2^T + A_3^T^2$$

(vii) 
$$A_1 = -1015.07 + 79.5387 \times -2.358016 \times^2 + 0.03031583 \times^3 - (1.400261 E - 4) \times^4$$

(viii) 
$$A_2 = 4.68103 - (3.037766 E - 1) x + (8.44845 E - 3)x^2$$
  
-  $(1.047721 E - 4) x^3 + (4.80097 E - 7)x^4$ 

(ix) 
$$A_3 = -(4.9107 E - 3) + (3.83184 E - 4) x$$
  
 $-(1.078963 E - 5) x^2 + (1.3152 E - 7) x^3$   
 $-(5.897 E - 10) x^4$ 

T = Solution temperature, °F, having range 60°F  $\geq$  T  $\leq$  330°F × = LiBr concentration, percent, having range 40% $\geq$ x $\leq$  70% The properties listed below are in S.I. units (h, kJ/kg; C<sub>p</sub>, kJ/kg; T, °C)

- (x) Specific heat of LiBr-water solution:  $C_{p_L} = 4.259 0.053343 \times + (2.307 E 4)x^2$
- (xi) Enthalpy of saturated water:  $h_{f} = (0.99615 T + (1.8239 E - 6)T^{2} - 0.13468 * (10)$  \*\* (-0.036 T)) + 0.13468) \* 4.187
- (xii) Enthalpy of saturated vapour:  $h_g = H_{fg} + h_f$
- (xiii) Latent heat of vaporization:  $h_{fg} = (597.31 - 0.555 T - 0.2389 * (10 * * (5.1463 - 1540/(T + 273.15)))) * 4.187$
- (xiv) Enthalpy of superheated vapour:  $h_{sp} = 1.925 * T_{g} - 0.126 T_{c} + 2500$
- (xv) Specific heat of water:  $C_{pw} = 4.2097187 - (1.4125 E - 3) T + (1.375 E - 5)T^2$

# COMPUTER RESULTS FOR OPTIMISATION OF GENERATOR TEMPERATURE TO

# CALCULATION OF PROPERTY VALUES AT GENERATOR TEMPERATURE PG=80 MM OF MERCURY

TC =	47.078	TA= 47.078	TE= 5,00004	X1=61.38680
41 =	135.785	H6 = 2510.082	H60=2510.082	H4 = 196.926
H40=	196.926	E1 = 0.000	$\begin{array}{c} \text{TE} = 5.00004 \\ \text{H6D} = 2510.082 \\ \text{E2} = 0.750 \end{array}$	PG= 80.000

The state of the s	1 X I	117 T	H70	ī	112D	1. Н3	T COP I
1 100.605 102.363 3 104.151 105.970 107.899 109.699 1113.5527 10 1117.53	153.700 154.400 165.100 165.800	244.860 251.279 257.800 264.413 271.107 277.871 284.694 291.565 298.474 305.409	146,124 150,363 154,692 159,093 163,547 168,034 172,533 177,021 181,474 185,867	Annual Wilder States Annual An	208 941 209 9424 210 499 211 275 212 829 213 458 215 204	2687.733 2691.117 2694.559 2698.060 2701.619 2705.238 2708.917 2712.656 2716.457 2720.319	0.60151 0.64538 0.67092 0.68734 0.69860 0.70668 0.71268 0.71720 0.72085 0.72374

## CALCULATION OF COEFFICIENTS OF EQ. (2.16)

A2 A4 A6 A8	= -0.537664E-06 = 0.493741E-10	A3= A5= A7= A9=	0.676124E-04 0.852616E-09 -0.662301E-12 -0.246315E-17
	Err \$ age	CDP 0.601506 0.645381 0.670923 0.687345 0.698680 0.7126680 0.7126680 0.712641 0.720853	

# VALUES CORRESPONDING TO OPTIMUM GENERATOR TEMPERATURE TG

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6.00	Co. Co.	1.30
Y	455 16	3
776 553 329	CT CR C1 455.16 526.0250 0.2 4082.2 455.16 814.7799 0.4 6659.3 664.82 1053.9344 0.6 8865.5 829.76	2
. 253	, 14.4477 X.2 8855.5 829.76	3
. 3 2 9	1493.7347	4

COP= 0.727325 QGAS=15412.98

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